WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

C08K 5/09

(11) International Publication Number:

WO 99/48963

A2

(43) International Publication Date: 30 September 1999 (30.09.99)

(21) International Application Number:

PCT/US99/06379

(22) International Filing Date:

24 March 1999 (24.03.99)

(30) Priority Data:

330077 09/127.316 25 March 1998 (25.03.98) 31 July 1998 (31.07.98)

NZ US

(71) Applicants: CHEVRON CHEMICAL COMPANY LLC [US/US]; 555 Market Street, San Francisco, CA 94105 (US). SEALED AIR (NZ) LIMITED [NZ/NZ]; Private Bag 3085, Corner Avalon Drive & Foreman Road (NZ).

(72) Inventors: CHING, Ta, Yen; 10 Santa Yorma Court, Novato, CA 94945 (US). CAI, Gangfeng; 335 Glasgow Circle, Danville, CA 94526 (US). DEPREE, Craig; 27 Pembroke Street, Palmerston North (NZ). GALLAND, Mark, Steven; 2806 Chasse Ridge Drive, Orange, TX 77632 (US). GOODRICH, Joseph, L.; 3545 Wilkinson Lane, Lafayette, CA 94549 (US). LEONARD, James, P.; 545 Bret Harte Road, San Rafael, CA 94901 (US). MATTHEWS, Andrew; 106 West Shallowstone Road, Greer, SC 29650 (US). RUSSELL, Kenneth, W.; Route 10, Box 1366, Orange, TX 77630 (US). YANG, Hu; 360 Bellflower Court, San Ramon, CA 94583 (US).

(74) Agents: HAYMOND, W., Bradley et al.; Chevron Corporation. Law Dept., P.O. Box 7141, San Francisco, CA 94120-7141

(81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

Without international search report and to be republished upon receipt of that report.

(54) Title: OXYGEN SCAVENGERS WITH REDUCED OXIDATION PRODUCTS FOR USE IN PLASTIC FILMS AND BEVERAGE AND FOOD CONTAINERS

FMCM poly(ethylene/methyl acrylate/cyclohexene-methyl acrylate)

(57) Abstract

An oxygen scavenger composition, for use in or with plastics materials, includes a polymer or oligomer having at least one cyclohexene group or functionality. The composition produces only low levels of volatile or (extractable from a plastics material in which it is incorporated) products as a consequence of oxygen scavenging. A family of polymers containing selected cyclic allylic pendent groups for oxygen scavenging packaging which has minimal organoleptic by-products after oxidation. Multilayer plastic containers for food and beverage packaging which incorporate oxygen scavenging resins which selectively oxidize upon activation without giving off odorous fragments.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑÜ	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	ĬL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	rs	Iceland	MW	Malawi	US	United States of Americ
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

1	OXYGEN SCAVENGERS WITH REDUCED OXIDATION PRODUCTS FOR
2	USE IN PLASTIC FILMS AND BEVERAGE AND FOOD CONTAINERS
3	
4	
5	Field of the Invention
6	The present invention is directed to oxygen scavengers for use in plastics
7	materials, and in particular plastics films. Emphasis is given to scavengers
8	which produce low or negligible levels of oxidation by-products which may
9	contaminate the head space in a package. This invention also relates to a
10	composition useful in scavenging oxygen from environments containing
11	oxygen-sensitive products, particularly food and beverage products. More
12	specifically, the oxygen scavenging composition includes a polymer having
13	ethylenic unsaturation contained within a cyclic moiety, a transition metal
14	compound and, optionally, a photoinitiator. The present invention also relates
15	to compositions for use in areas such as food packaging, and with minimal
16	effect on odor and taste of packaged contents. The invention preferably uses .
17	ethylene acrylate copolymers which are modified with selected cyclic allylic
18	pendent groups for use in oxygen scavenging packaging materials. The
19	present invention also relates to rigid polymeric food or beverage containers
20	comprising polyester such as polyester terephthalate or polyester naphthalate
21	and oxygen scavenging polymer.
22	
23	
24	
25	Background of the Invention
26	
27	The majority of plastic films produced are employed in some form of
28	packaging. The present invention is primarily concerned with those films
29	used for applications requiring a low level of oxygen in a package, though
30	may also find other uses.

1	Limiting the exposure of oxygen-sensitive products to oxygen maintains and
2	enhances the quality and shelf life of many products. For instance, by limiting
3	the oxygen exposure of oxygen-sensitive food products in a packaging
4	system, the quality of the food product can be maintained and spoilage
5	retarded. In addition, such packaging also keeps the product in inventory
6	longer, thereby reducing costs incurred from waste and having to restock.
7	
8	In the food packaging industry, several techniques for limiting oxygen
9	exposure have been developed. Common techniques include those where
10	oxygen is consumed within the packaging environment by some means other
11	than the packaged article or the packaging material (e.g., through the use of
12	oxygen scavenging sachets), those where reduced oxygen environments are
13	created in the package (e.g., modified atmosphere packaging (MAP) and
14	vacuum packaging), and those where oxygen is prevented from entering the
15	packaging environment (e.g., barrier films).
16	
17	The art dealing with barrier packaging, and the low oxygen or modified
18	packaging of products is relatively well developed. This includes the use of
19	films and inserts containing oxygen scavenging compounds able to extract a
20	majority of any residual oxygen after packaging occurs.
21	
22	Oxygen scavenging compounds for use in plastic films are relatively well
23	known. Typically these comprise unsaturated compounds in combination with
24	a transition metal catalyst. In response to some form of initiation—usually
25	exposure to light or radiation—the scavengers react with available oxygen in
26	the package.
27	
28	For instance:
29	

Michael Rooney, "Oxygen scavenging: a novel use of rubber photo-1 2 oxidation", Chemistry and Industry, March 20, 1982, pp. 197-198, describes the use of ethylenically unsaturated compounds as oxygen scavengers on 3 exposure to light. However, systems describing the use of transition metal 4 5 catalysts are not described. 6 US 4,908,151 to Mitsubishi describes sachets containing unsaturated fatty 7 acid (i.e., an ethylenically unsaturated hydrocarbon) in combination with a 8 9 transition metal compound in a basic substance. However, there is no 10 description of these materials in the form of a film nor the use of photo-11 exposure as an initiating mechanism. 12 Japanese patent JP5032277 to Kuwa describes the use of radical containing 13 14 resin layers in packages. The invention comprises an oxidizable polymer 15 whose oxygen scavenging abilities is photoinitiated. 16 New Zealand patent application NZ241802 to W R Grace and also NZ243077 17 18 also to WR Grace, claim oxygen scavenging compositions comprising 19 ethylenically unsaturated hydrocarbons with transition metal catalysts. A wide 20 range of ethylenically unsaturated compounds are discussed in the texts of 21 these specifications though there is no mention of the problems to which the 22 present invention is directed, nor the compounds and products encompassed 23 by the present invention. 24 25 Sachets containing an oxygen scavenging compositions can contain ferrous 26 compositions, which oxidize to their ferric state, unsaturated fatty acid salts on 27 an absorbent, and/or a metal-polyamide complex. See, e.g., U.S. Patent 28 Nos. 4,908,151 and 5,194,478. The disadvantages of sachets include the 29 need for additional packaging steps (to add the sachet to the package), the

-4-

1 potential for contamination of the packaged article should the sachet break 2 and the danger of ingestion by a consumer. 3 4 Oxygen scavenging materials also have been incorporated directly into the 5 packaging structure. This technique (hereinafter referred to as "active oxygen 6 barrier") can provide a uniform scavenging effect throughout the package and 7 can provide a means of intercepting and scavenging oxygen as it passes 8 through the walls of a package, thereby maintaining the lowest possible 9 oxygen level throughout the package. Active oxygen barriers have been 10 formed by incorporating inorganic powders and/or salts as part of the 11 package. See, e.g., U.S. Patent Nos. 5,153,038, 5,116,660, 5,143,769, and 12 5,089,323. However, incorporation of such powders and/or salts can degrade 13 the transparency and mechanical properties (e.g., tear strength) of the 14 packaging material and can complicate processing, especially where thin 15 films are desired. Also, these compounds as well as their oxidation products 16 can be absorbed by food in the container, which can result in the food product 17 failing to meet governmental standards for human consumption. 18 19 EP 0 519 616 discloses an oxygen scavenging composition that includes a 20 blend of an epoxide, a first polymeric component grafted with an unsaturated 21 carboxylic anhydride and/or acid, a second polymeric component including OH, SH, or NHR² groups where R² is H, C₁-C₃ alkyl, or substituted C₁-C₃ alkyl 22 23 moiety, and a metal salt capable of catalyzing the reaction between oxygen and the second polymeric component. The first polymeric component is 24 present in an amount sufficient to ensure that the blend is non-phase 25 26 separated. A blend of polymers is utilized to obtain oxygen scavenging, and 27 the second polymeric component is preferably a (co)polyamide such as 28 MXD6. 29

29

1 Another type of active oxygen barrier is illustrated in EP-A-0 301 719, 2 EP-A-0 380 319, PCT Publication No. WO 90/00578, and PCT Publication 3 No. WO 90/00504. See also U.S. Patent Nos. 5,021,515 5,194,478, and 4 5,159,005. The disclosed oxygen scavenger includes polyamide-transition 5 metal catalyst compositions. Through catalyzed scavenging by the 6 polyamide, the package wall regulates the amount of oxygen reaching the 7 interior of the package. However, the onset of useful oxygen scavenging (i.e., up to about 5.8 x 10⁻⁵ cm³/m²•24 hours at ambient conditions) can take as 8 9 long as 30 days to occur. Therefore, this technique is not acceptable for 10 many applications. Further, polyamides typically are incompatible with many 11 thermoplastic polymers commonly used to make flexible packaging materials 12 (e.g., ethylene/vinyl acetate copolymers, low density polyethylene, etc.) or, 13 when used by themselves, are difficult to process and result in inappropriately 14 stiff structures. 15 16 Oxygen scavenging compositions that include transition metal catalysts and 17 ethylenically unsaturated hydrocarbon polymers which have an ethylenic 18 double bond content of from 0.01 to 10 equivalents per 100 grams of polymer 19 are disclosed in U.S. Patent No. 5,399,289. Various conventional 20 homopolymers, copolymers, and polymer blends are disclosed. Because 21 these polymers are amorphous, they can be difficult to blend and process 22 with film-forming semicrystalline polymers conventionally used to make 23 flexible packaging materials. 24 25 The use of a transition metal and a photoinitiator to facilitate initiation of 26 effective scavenging activity of ethylenically unsaturated compounds is taught 27 in U.S. Patent No. 5,211, 875, which is incorporated herein by reference as if 28 set forth in full.

1 PCT Publication Nos. WO 95/02616 and WO 96/40799 disclose a scavenger 2 composition that includes a transition metal salt and a copolymer (of ethylene 3 and a vinyl monomer) having ether, amino, carboxylic acid, ester, or amide functionalities pendent therefrom. Although these compositions can provide 4 5 oxygen scavenging activity, the particular advantages of having ethylenic 6 unsaturation contained within a cyclic mojety are not disclosed. Because the 7 compositions of this invention are significantly cleaner than those described in 8 the prior art, they do not require the use of high levels of adjuncts to absorb 9 the undesirable byproducts. Such absorbent additives are known in the art. 10 for example see U.S. 5,834,079 and U.S. 08/857,276. It is also well known in 11 the art that such additives (zeolites and silicas) adversely affect the haze and 12 clarity of packaging structures. 13 14 PCT Application WO 96/40799 from Chevron describes the use of a variety of 15 ethylenic materials with benzylic, allylic or ether containing side chains. Some 16 of these materials may be prepared by esterification or transesterification of a 17 polymer melt. The use of pendent cyclic groups containing allylic 18 unsaturation is generally referred to, but there is only one such example, 19 wherein Nopol, a bicyclic alcohol, is used in a transesterification reaction and 20 oxygen absorbing films are formulated from the product. There is no 21 reference to the benefits of cyclic allylic compounds as described in this 22 invention i.e., on oxidation they produce very low levels of oxidation 23 byproducts when compared to comparable linear allylic systems. Because of 24 its bicyclic nature. Nopol is not expected to produce these benefits. 25 26 While the prior art compounds may effectively scavenge oxygen they 27 introduce other problems into packaging. For instance, in summary the prior 28 art incorporates into film structures compounds which are ethylenically 29 unsaturated but which often cleave as a consequence of the reactions of the 30 oxygen scavenging process. For example, films containing unsaturated

WO 99/48963

1	compounds such as squalene or vegetable oils produce large amounts of
2	volatile aldehydes and ketones upon oxidation. Unfortunately, many of these
3	volatile compounds are not contained within the film structure and find their
4	way into the head space of the package. Here they can represent more of a
5	problem than the oxygen which they have replaced and have the potential to
6	contaminate comestible products.
7	
8	This problem represents a significant problem yet has been downplayed or
9	overlooked by the published prior art. As a consequence, those searching
10	the prior art for a solution to this problem find no answer—the art appears to
11	be directed primarily along a narrow track of improving on scavenging
12	efficiencies, or physical properties of scavenging films, rather than
13	recognizing or addressing other associated problems.
14	
15	Accordingly the present invention seeks to address the problems associated
16	with scission products of oxygen scavengers, and seeks also to provide a
17	group of compounds and substances (as well as films and plastics materials
18	including same) which have an advantage over the prior art in terms of
19	reduced quantities of scission products.
20	
21	Ideally, a polymeric material for use in an oxygen scavenging composition
22	should exhibit good processing characteristics, be able to be formed into
23	useful packaging materials or have high compatibility with those polymers
24	commonly used to make packaging materials, and not produce byproducts
25	which detract from the color, taste, or odor of the packaged product. It has
26	been found that when the ethylenic unsaturation is contained within a cyclic
27	group, substantially fewer and less byproducts are produced upon oxidation
28	as compared to analogous non-cyclic materials. Optimally, a packaging

1 material formed from such a composition can retain its physical properties 2 after significant oxygen scavenging. 3 4 5 New polymer compositions having properties that are particularly tailored for 6 specific applications are required in response to more specific and 7 sophisticated end uses. It can be difficult to make these compositions directly 8 by polymerization from monomers or via solution esterification or 9 transesterification, but manufacturing them in melt mixing equipment such as 10 an extruder has provided an efficient, economical and viable means to supply 11 increasingly complex polymers to meet the needs in specialized markets. 12 It is well known that regulating the exposure of oxygen-sensitive products to 13 oxygen maintains and enhances the quality and "shelf-life" of the product. 14 For instance, by limiting the exposure of oxygen sensitive food products to 15 oxygen in a packaging system, the quality or freshness of food is maintained. 16 spoilage reduced and the food shelf life extended. In the food packaging 17 industry, several means for regulating oxygen exposure have already been 18 developed. These means include modified atmosphere packaging (MAP) and 19 oxygen barrier film packaging. 20 One method currently being used is "active packaging", whereby the package 21 containing the food product has been modified in some manner to regulate 22 the food's exposure to oxygen. One form of active packaging uses oxygen-23 scavenging sachets which contain a composition which scavenges the 24 oxygen through oxidation reactions. One type of sachet contains iron-based 25 compositions which oxidize to their ferric states. Another type of sachet 26 contains unsaturated fatty acid salts on a particulate adsorbent. Yet another 27 sachet contains metal/polyamide complex. However, one disadvantage of

- 1 sachets is the need for additional packaging operations to add the sachet to
- 2 each package. A further disadvantage arising from the iron-based sachets is
- 3 that certain atmospheric conditions (e.g., high humidity, low CO₂ level) in the
- 4 package are sometimes required in order for scavenging to occur at an
- 5 adequate rate. Further, the sachets can present a problem to consumers if
- 6 accidentally ingested.
- 7 Another means for regulating exposure of a packaged product to oxygen
- 8 involves incorporating an oxygen scavenger into the packaging structure
- 9 itself. A more uniform scavenging effect through the package is achieved by
- 10 incorporating the scavenging material in the package instead of adding a
- 11 separate scavenger structure (e.g., a sachet) to the package. This may be
- 12 especially important where there is restricted airflow inside the package. In
- addition, incorporating the oxygen scavenger into the package structure
- 14 provides a means of intercepting and scavenging oxygen as it permeates the
- walls of the package (herein referred to as an "active oxygen barrier"),
- thereby maintaining the lowest possible oxygen level in the package.
- 17 One attempt to prepare an oxygen-scavenging wall involves the incorporation
- 18 of inorganic powders and/or salts. However, incorporation of these powders
- 19 and/or salts causes reduction of the wall's optical transparency, discoloration
- 20 after oxidation, and reduced mechanical properties such as tear strength. In
- 21 addition, these compounds can lead to processing difficulties, especially
- 22 when fabricating thin films. The oxidation products may migrate into food at
- 23 levels which would not be regarded as safe or can impart unacceptable taste
- 24 or smell to food.
- 25 An oxygen-scavenging composition comprising a blend of a first polymeric
- component comprising a polyolefin is known, the first polymeric component
- 27 having been grafted with an unsaturated carboxylic anhydride or an
- 28 unsaturated carboxylic acid, or combinations thereof, or with an epoxide; a

- 1 second polymeric component having -OH, -SH, or -NHR² groups where R² is
- 2 H, C_1 - C_3 alkyl, substituted C_1 - C_3 alkyl; and a catalytical amount of metal salt
- 3 capable of catalyzing the reaction between oxygen and the second polymeric
- 4 component, the polyolefin being present in an amount sufficient so that the
- 5 blend is not phase-separated. A blend of polymers is utilized to obtain
- 6 oxygen scavenging, and the second polymeric component is preferably a
- 7 polyamide or a copolyamide such as the copolymer of m-xylylene-diamine
- 8 and adipic acid (MXD6).
- 9 Some oxygen scavenging systems produce an oxygen-scavenging wall. This
- 10 is done by incorporating a metal catalyst-polyamide oxygen scavenging
- 11 system into the package wall. Through catalyzed oxidation of the polyamide,
- 12 the package wall regulates the amount of oxygen which reaches the interior
- 13 volume of the package (active oxygen barrier) and has been reported to have
- 14 oxygen scavenging rate capabilities up to about 5 cubic centimeters (cc)
- 15 oxygen per square meter per day at ambient conditions. However, this
- 16 system suffers from significant disadvantages.
- 17 One particularly limiting disadvantage of polyamide/catalyst materials can be
- 18 a low oxygen scavenging rate. Adding these materials to a high-barrier
- 19 package containing air can produce a package which is not generally suitable
- 20 for creating an internal oxygen level of less than 0.1% within seven days at
- 21 storage temperatures, as is typically required for headspace oxygen
- 22 scavenging applications.
- 23 There are also disadvantages to having the oxygen-scavenging groups in the
- 24 backbone or network structure in this type of polyamide polymer. The basic
- 25 polymer structure can be degraded and weakened upon reaction with oxygen.
- 26 This can adversely affect physical properties such as tensile or impact
- 27 strength of the polymer. The degradation of the backbone or network of the

- 1 polymer can further increase the permeability of the polymer to those
- 2 materials sought to be excluded, such as oxygen.
- 3 Moreover, polyamides previously used in oxygen scavenging materials, such
- 4 as MXD6, are typically incompatible with thermoplastic polymers used in most
- 5 flexible packaging walls, such as ethylene-vinyl acetate copolymers and low
- 6 density polyethylene. Even further, when such polyamides are used by
- 7 themselves to make a flexible package wall, they may result in inappropriately
- 8 stiff structures. They also incur processing difficulties and higher costs when
- 9 compared with the costs of thermoplastic polymers typically used to make
- 10 flexible packaging. Even further, they are difficult to heat seal. Thus, all of
- 11 these are factors to consider when selecting materials for packages,
- 12 especially multi-layer flexible packages and when selecting systems for
- 13 reducing oxygen exposure of packaged products.
- 14 Another approach to scavenging oxygen is an oxygen-scavenging
- 15 composition comprising an ethylenically unsaturated hydrocarbon and a
- transition metal catalyst. Ethylenically unsaturated compounds such as
- 17 squalene, dehydrated castor oil, and 1,2-polybutadiene are useful oxygen
- 18 scavenging compositions, and ethylenically saturated compounds such as
- 19 polyethylene and ethylene copolymers are used as diluents. Compositions
- 20 utilizing squalene, castor oil, or other such unsaturated hydrocarbon typically
- 21 have an oily texture as the compound migrates toward the surface of the
- 22 material. Further, polymer chains which are ethylenically unsaturated in the
- 23 backbone would be expected to degrade upon scavenging oxygen,
- 24 weakening the polymer due to polymer backbone breakage, and generating a
- 25 variety of off-odor/off-taste by-products.
- 26 Other oxidizable polymers recognized in the art include "highly active"
- 27 oxidizable polymers such as poly(ethylene-methyl acrylate-benzyl acrylate),
- 28 EMBZ, and poly(ethylene-methyl acrylate-tetrahydrofurfuryl acrylate), EMTF,

28

1 as well as poly(ethylene-methyl acrylate-nopol acrylate), EMNP. Although effective as oxygen scavengers, these polymers have the drawback of giving 2 3 off large amounts of volatile by-products and/or strong odors after oxygen 4 scavenging. Also known are oxygen-scavenging compositions which comprise a transition-5 6 metal salt and a compound having an ethylenic backbone and having allylic 7 pendent or terminal moieties which contain a carbon atom that can form a 8 free radical that is resonance-stabilized by an adjacent group. Such a 9 polymer needs to contain a sufficient amount and type of transition metal salt 10 to promote oxygen scavenging by the polymer when the polymer is exposed 11 to an oxygen-containing fluid such as air. Although effective as oxygen 12 scavengers, upon oxidation, we have found that allylic pendent groups on an 13 ethylenic backbone tend to generate considerable amounts of organic 14 fragments. We believe this is a result of oxidative cleavage. We believe 15 these fragments can interfere with the use of allylic pendent groups as 16 oxygen scavengers in food packaging. 17 Multilayer rigid container structures, which utilize an oxygen scavenging 18 composition, are known. In the container wall, base polymers such as polyethylene terephthalate have been used along with an oxygen scavenger. 19 20 The resulting multilayer package wall includes at least an oxygen scavenger 21 core layer as well as inner and outer layers having high oxygen barrier 22 qualities. The oxygen scavenger core layer is a combination of at least an 23 oxygen scavenging polymer with post consumer-polyethylene terephthalate 24 (PC-PET). The inner and outer layers include at least oxygen barrier quality 25 PET. 26 27 Furthermore, multilayered plastic bottles having oxygen scavenging capacity

sufficient to maintain substantially zero or near zero presence of oxygen in

1 the bottle cavity under specified storage conditions have also been disclosed. 2 The multilayered bottle wall has at least three layers. The inner and outer 3 layers are PET or another bottling polyester, which define the bottle cavity 4 and the outside skin of the bottle respectively. Between the inner and outer 5 layers is an oxygen scavenging copolyester layer. 6 7 Condensation copolymers used for making bottles with polyester such as 8 PET or polyethylene naphthalate (PEN) have also been disclosed. The 9 condensation copolymers comprise predominantly polyester segments and 10 an oxygen scavenging amount of polyolefin oligomer segments. The 11 copolymers are preferably formed by transesterification during reactive 12 extrusion and typically comprise about 0.5 to about 12 wt. % of polyolefin 13 oligomer segments. In a preferred embodiment, a bottle is provided having a 14 multilayer wall of at least three layers. The outer and inner layers are of 15 unmodified PET and the oxygen scavenging layer in between the outer and 16 inner layer is made of the condensation copolymers described above having 17 an oxygen scavenging amount of polyolefin oligomers. 18 19 A transparent oxygen-scavenging article for packaging oxygen sensitive 20 products is also known, the oxygen-scavenging article having a multilayered 21 wall including at least three layers, an inner and outer layer of biaxially-22 oriented aromatic polyester polymers such as PET or PEN and an oxygen-23 scavenging aromatic ester polymer compatible with the polyester polymer. 24 The oxygen-scavenging aromatic ester polymer must include ketone carbonyl groups to provide the oxygen-scavenging functionality and aromatic and ester 25 26 groups for compatibility with the polyester. 27 28 PET containers have been disclosed that have a container wall of stretched 29 plastic material with high oxygen barrier properties and an activating metal 30 incorporated into the plastic material. The plastic material is PET in admixture

30

with a polyamide and the metal is either added to the mixture or contained in 1 2 one or both of the polymers. 3 4 A container containing at least one layer containing a plastics material and 5 ions of at least one metal has also been disclosed. The plastics material in the layer consists of at least a partially split or degraded polyamide which has 6 7 increased sensitivity to reaction with oxygen in the presence of metal thus 8 giving the layer improved oxygen barrier properties. 9 A container has been disclosed with a wall having high oxygen barrier 10 11 properties comprising a molded polymer composition, the composition 12 comprising a granular mixture of (1) a first polymer providing essential strength for the container wall and (2) an active component comprising a 13 14 metal compound capable of scavenging oxygen and consisting essentially of 15 a metal ion having complexing properties and a polymer to which said metal ion is combined as a metal complex in the molded polymer composition of 16 said wall to scavenge oxygen. There is also disclosed a method of producing 17 the polymer composition which can be molded into containers, the method 18 being to treat a polymer with a metal compound dissolved or slurried in a 19 volatile solvent composition during refluxing conditions for obtaining the active 20 21 component having capacity to scavenge oxygen. 22 23 An article has been disclosed with oxygen barrier properties comprising at least partly a molded polymer composition formed by melting granules of the 24 25 composition and molding the melted composition to produce the article. The 26 composition comprises a granular mixture of (1) a first polymer composition providing strength for the article and (2) a second polymer composition 27 28 compatible with the first polymer composition. The second polymer 29 composition is obtainable by reacting a polyamide or copolyamide with a

solution of a transition metal compound in a volatile solvent under refluxing

1 conditions. The polymer of the first polymer composition can be any polymer 2 and the metal of the metal compound reacted with the polyamide or 3 copolyamide can be any transition metal. The amount of metal in the second 4 polymer composition is at least 500 ppm. 5 6 A polymer material having increased sensitivity to reaction with oxygen has 7 also been disclosed, the polymer material comprising a polyamide, which has 8 been reacted with a nucleophilic reagent and possibly an activator. The 9 nucleophilic reagent is selected from the group consisting of compounds containing at least one hydroxyl group, compounds containing at least one 10 11 alkoxide group, phosphate compounds, pyrophosphate compounds. 12 polyphosphate compounds, salts of organic acids and a copolymer of vinyl 13 alcohol and ethylene. The activator is in the form of a hydrogen donor. A 14 process is also disclosed for producing the polymer material, which has increased sensitivity of reaction with oxygen. In the process, a polyamide 15 16 reacts with the nucleophilic reagent under such conditions that the polymer 17 material is obtained. 18 19 Such polymeric containers of PET, PEN and/or polyamide as described 20 above utilize oxidizable components to react with and decrease the amount of 21 oxygen in contact with oxygen sensitive materials packaged in containers. All 22 of these oxidizable materials have the disadvantage of imparting unpleasant 23 odor and/or taste to the packaged materials because of the byproducts given off during the oxidation of the oxidizable materials. Another problem is the 24 uncontrolled oxidation fragmentation from the polymer backbone which leads 25 26 to chain secession, thus weakening the physical integrity of the multilayer 27 container structures. 28

29

-16-

1 The present invention solves many of the problems of the prior art, especially 2 with an oxygen scavenging packaging material incorporating polymers 3 comprising cyclic allylic (olefinic) pendent groups which produce little or no 4 migration of oxidation by-products adversely affecting odor or taste, thus 5 minimizing organoleptic problems in food packaging. This is because the 6 cyclic allylic structures are less likely to fragment or cleave after oxidation than the conventional open chain allylic (olefinic) groups used in oxygen 7 8 scavenging packaging material. 9 10 Such polymeric containers of PET, PEN and/or polyamide as described 11 above utilize oxidizable components to react with and decrease the amount of 12 oxygen in contact with oxygen sensitive materials packaged in containers. All 13 of these oxidizable materials have the disadvantage of imparting unpleasant 14 odor and/or taste to the packaged materials because of the byproducts given 15 off during the oxidation of the oxidizable materials. Another problem is the 16 uncontrolled oxidation fragmentation from the polymer backbone which leads 17 to chain secession, thus weakening the physical integrity of the multilayer 18 container structures. 19 20 In contrast, the present invention achieves a rigid beverage and food 21 container comprising PET and/or PEN, the container incorporating an oxygen 22 scavenging component of cyclic olefin which does not give off odor and or 23 taste as a result of its oxygen scavenging function. The oxidation also does 24 not cause a change in molecular weight. This is because the cyclic olefin 25 oxygen scavenging component does not fragment as it oxidizes, thus 26 avoiding the problem of imparting oxidation byproducts to the packaged

material while maintaining the structural integrity.

27

28

1	It is an object of the present invention to address the foregoing problems or at
2	least to provide the public with a useful choice.
3	
4	Further aspects and advantages of the present invention will become
5	apparent from the ensuing description, which is given by way of example
6	only.
7	
8	SUMMARY OF INVENTION
9	
10	According to one aspect of the present invention, there is provided an oxygen
11	scavenger for use in or with plastics materials, said scavenger comprising or
12	including a polymer or oligomer having at least one cyclohexene group or
13	functionality.
14	,
15	According to another aspect of the present invention, there is provided an
16	oxygen scavenger, substantially as described above, which produces only
17	low levels of volatile or extractable (from a plastics material in which it is
18	incorporated) products as a consequence of oxygen scavenging.
19	
20	According to another aspect of the present invention there is provided an
21	oxygen scavenger, substantially as described above, which is substantially
22	stable with respect to reaction with oxygen until triggered by an external
23	event.
24	
25	According to another aspect of the present invention, there is provided an
26	oxygen scavenger, substantially as described above, wherein the external
27	event is irradiation by electromagnetic radiation.
28	,
29	According to a further aspect of the present invention, there is provided an
30	oxygen scavenging composition, including an oxygen scavenger,

а

1	substantially as described above, which includes one of more (rigger
2	enhancing components making the scavenger susceptible to triggering from
3	an external event.
4	
5	According to another aspect of the present invention, there is provided an
6	oxygen scavenging composition, substantially as described above, wherein a
7	trigger-enhancing component may be benzophenone or substituted
8	derivatives thereof.
9	
10	According to another aspect of the present invention, there is provided an
11	oxygen scavenging composition, substantially as described above, which
12	includes the presence of one or more catalysts for the scavenging process.
13	
14	According to another aspect of the present invention, there is provided an
15	oxygen scavenging composition, substantially as described above, in which a
16	catalyst may be a transition metal salt, compound or complex.
17	
18	According to another aspect of the present invention, there is provided an
19	oxygen scavenger or oxygen scavenging composition, substantially as
20	described above, which is in the form of a plastics resin.
21	
22	According to another aspect of the present invention, there is provided an
23	oxygen scavenger or oxygen scavenging composition, substantially as
24	described above, in which the plastics resin is a resin suitable for use in the
25	manufacture of plastic films.
26	
27	According to another aspect of the present invention, there is provided an
28	oxygen scavenger or oxygen scavenging composition, substantially as
29	described above, in which the plastic resin is a polyester resin.
30	

1	According to another aspect of the present invention, there is provided an
2	oxygen scavenger or oxygen scavenging composition, substantially as
3	described above, when present in a plastics film or layer thereof.
4	
5	According to another aspect of the present invention, there is provided an
6	oxygen scavenger or oxygen scavenging composition, substantially as
7	described above, when used as a polymeric material of a plastics film, a layer
8	thereof, and/or a coating thereof, or in a plastics material.
9	
10	According to another aspect of the present invention, there is provided an
11	oxygen scavenger or oxygen scavenging composition, substantially as
12	described above, when dispersed throughout a plastics film, a layer thereof,
13	and/or a coating thereon, or in a plastics material.
14	· ·
15	According to a further aspect of the present invention, there is provided an
16	oxygen scavenger or oxygen scavenging composition, substantially as
17	described above, in which the anhydride comprises 1,2,3,6-tetrahydrophthalic
18	anhydride or tetrahydrophthalic anhydride monomer derivable from
19	butadiene.
20	
21	According to a further aspect of the present invention, there is provided an
22	oxygen scavenger or oxygen scavenging composition prepared from the
23	reaction of a tetrahydrobenzyl alcohol with one or more compounds having
24	one or more of the following functionalities: carboxylic acid, acid halide, ester,
25	anhydride, and isocyanate.
26	
27	According to another aspect of the present invention, there is provided an
28	oxygen scavenger or oxygen scavenging composition, substantially as
29	described above, in which the alcohol comprises tetrahydrobenzyl alcohol.
30	

1 According to another aspect of the present invention, there is provided an 2 oxygen scavenger or oxygen scavenging composition, substantially as 3 described above, in the compounds with which the alcohol is reacted may 4 include a styrene maleic anhydride copolymer, and/or a polyfunctional 5 isocvanate. 6 7 According to another aspect of the present invention, there is provided an 8 oxygen scavenger or oxygen scavenging composition, prepared from a 9 cyclohexene dimethanol compound. 10 11 According to another aspect of the present invention, there is provided an 12 oxygen scavenging polymer including at least one pendant cyclohexene 13 group prepared by a reactive extrusion process. 14 15 According to a further aspect of the present invention, there is provided an 16 oxygen scavenger or oxygen scavenging polymer, substantially as described 17 above, in which the reactive extrusion process comprises an esterification or 18 transesterification step. Suitable catalyst include acids, bases and 19 organometallic compounds such as the titanium alkoxides. 20 21 According to another aspect of the present invention, there is provided an 22 oxygen scavenger or oxygen scavenging polymer prepared by a route 23 including a cyclohexene anhydride. 24 25 According to another aspect of the present invention, there is provided an 26 oxygen scavenger or oxygen scavenging polymer prepared by a route 27 including the reaction of a diene monomer, or hydroxy containing monomer, 28 with a cyclic anhydride. 29

1	According to another aspect of the present invention, there is provided an
2	oxygen scavenger or oxygen scavenging polymer, substantially as described
3	above, in which the cyclic anhydride is a maleic anhydride.
4	
5	According to a further aspect of the present invention, there is provided an
6	oxygen scavenger including a pendant cyclic alkene group prepared via a
7	method including a Diels Alder addition reaction.
8	
9	According to another aspect of the present invention, there is provided an
10	oxygen scavenger, substantially as described above, in which the preferred
11	dienes for use in the Diels Alder reaction is substituted and/or unsubstituted
12	1,3 butadiene.
	· · · · · · · · · · · · · · · · · · ·
13	According to another aspect of the present invention, there is provided an
14	oxygen scavenger, substantially as described above, in which the preferred
15	dienophile for use in the Diels Alder reaction include unsaturated acids,
16	anhydrides, and esters.
17	
18	According to another aspect of the present invention, there is provided an
19	oxygen scavenger, substantially as described above, in which the cyclic
20	alkene is cyclohexene.
21	
22	In other aspects, the present invention provides an article which include at
23	least one layer formed from a blend that includes the foregoing composition
24	as well as a method of scavenging oxygen in which a packaging article, at
25	least one layer of which is formed from a blend that includes the foregoing
26	composition, is exposed to actinic or e-beam radiation so as to activate the
27	composition.
28	

1 According to a further aspect of the present invention, there is provided an 2 oxygen scavenger or oxygen scavenging composition prepared from a 3 tetrahydrophthalic anhydride and a polymer or lower molecular weight 4 compound containing at least one amine group. 5 6 According to a further aspect of the present invention, there is provided an 7 oxygen scavenger or oxygen scavenging composition prepared from 8 diglcidyltetrahydrophthalate. 9 According to a further aspect of the present invention, there is provided an 10 oxygen scavenger or oxygen scavenging composition prepared from the 11 reaction of tetrahydrobenzyl alcohol, methyl or dimethyl substituted 12 tetrahydrobenzyl alcohol with one or more compounds having one or more of 13 the following functionalities: carboxylic acid, acid halide, ester, anhydride. 14 epoxide and isocyanate. 15 16 According to a further aspect of the present invention, there is provided an 17 oxygen scavenger or oxygen scavenging composition, substantially as 18 described above, in which a tetrahydrobenzyl alcohol or substituted 19 tetrahydrobenzyl alcohol reacts with one or more of the following materials: 20 21 ethylene (meth)acrylic acid and other acid containing polymers and acid 22 containing lower molecular weight materials: 23 24 styrene maleic anhydride copolymers; alpha olefin maleic anhydride 25 copolymers such as octadecene maleic anhydride; ethylene and ethylene 26 alpha olefin maleic anhydride terpolymers; ethylene alkyl (meth) acrylate 27 maleic anhydride terpolymers and other like anhydride containing polymers or 28 anhydride containing lower molecular weight materials:

29

7	polyment or lower molecular weight materials containing acid halide
2	functionality such as poly acryloyl chloride;
3	
4	ethylene alkyl (meth)acrylate copolymers and terpolymers and alternative
5	polymers or lower molecular weight materials containing lower alkyl ester
6	functionality;
7	•
8	epoxy resins,
9	
10	isocyanate functional material such as prepolymers and oligomers derived
11	from the common diisocyanates such as MDI, TDI and the like.
12	
13	According to a further aspect of the present invention, there is provided an
14	oxygen scavenger or oxygen scavenging composition prepared from a
15	dihydroxy cyclohexene compound. For example, 3 Cyclohexene-1,1-
16	dimethanol or its substituted derivatives may be used to prepare polyurethane
17	and polyester resins.
18	
19	According to a further aspect of the present invention, there is provided an
20	oxygen scavenger or oxygen absorbing composition prepared from a
21	cyclohexene carboxylic acid. Such materials may be prepared from acrylic
22	acid and substituted and unsubstituted butadienes. A typical example would
23	be tetrahydrobenzoic acid, derived from acrylic acid and butadiene. This may
24	be reacted with the following materials:
25	
26	hydroxyl functional materials such as poly(vinyl alcohol) and polyethylene-
27	vinyl alcohol, hydroxyl functional oligomers such as poly(ethylene glycol), the
28	polyester polyols and other lower molecular weight hydroxyl functional
29	materials;
30	

amine functional polymers and lower molecular weight compounds; 1 2 3 polyvalent metal ions. 4 5 According to a further aspect of the present invention, there is provided an 6 oxygen scavenger prepared from a cyclohexene functional acid chloride. 7 Example 9 utilizes 3-cyclohexene-1-carbonyl chloride. 8 9 According to a further aspect of the present invention, there is provided an 10 oxygen scavenger or oxygen scavenging composition prepared from 11 tetrahydrobenzaldehyde and its substituted derivatives. These may be 12 prepared from reaction of butadiene or the methyl substituted butadienes with 13 acrolein. 14 15 The tetrahydrobenzaldehydes may be reacted with hydroxyl functional 16 polymers such as poly(vinyl alcohol) and polyethylene-vinyl alcohol to form 17 polyvinyl acetals. 18 19 The following definitions apply herein throughout unless a contrary intention is 20 expressly indicated: 21 22 "polymer" means the polymerization product of one or more monomers and 23 includes homopolymers, as well as copolymers; 24 "copolymer" means the polymerization product of two or more kinds of 25 monomers; 26 27 "(meth)acrylate" means acrylate or methacrylate; 28 29 "photoinitiator" means a substance which, when activated by actinic radiation, 30 enhances and/or facilitates the initiation of one or more properties (e.g.,

1 oxygen scavenging) in another compound, thus resulting in a shorter 2 induction period and/or an increase in the rate of oxygen uptake of the overall 3 system: 4 5 "induction period" means the length of time beginning with the initiation of the 6 active components of a composition and ending with the onset of one or more 7 useful properties (e.g., oxygen scavenging); and 8 9 "antioxidant" means a material which can inhibit oxidative degradation and/or 10 crosslinking of a poly polymer so as to, for example, prolong the useful 11 lifetime of the polymer, to stabilize a polymer-containing composition during 12 processing (e.g., extrusion, coating, lamination, etc.); and/or to prolong the 13 shelf-life of the composition (prior to exposure thereof to actinic or e-beam 14 radiation). 15 16 The present invention is directed to oxygen scavengers. The invention 17 includes oxygen scavenging substances, as well as compositions containing 18 same. The form of the oxygen scavengers may vary and may comprise small 19 molecules through to large macromolecules as well as those sized in 20 between. The oxygen scavengers will be characterized in that they will be 21 able to react with oxygen in their near vicinity, enabling the removal of oxygen 22 from a closed system. 23 24 While the actual form of the oxygen scavengers may vary, a characteristic 25 that they each share is they include cyclic alkene groups or functionalities, 26 which are able to react with oxygen to provide the desired oxygen scavenging 27 properties. In preferred embodiments of the present invention, this will 28 comprise a cyclohexene group i.e. a six-membered ring with double bond 29 between two adjacent carbon atoms. It is acceptable that some carbons of

the cyclohexene group may also form a part of other ring structures within the 1 2 molecule, and/or form part of the skeleton of the molecule. It is not necessary 3 that the entire C₆ ring be dangling free of the remainder of the molecule to 4 which it is attached. A consideration however is that the group should be so 5 positioned and incorporated into the structure that the double bond is 6 available for reaction with oxygen. 7 8 It has been mentioned above that various scavengers of the present invention 9 may take different forms. This will also have some bearing on how they are 10 used, and also produced. Perhaps the simplest embodiments of the present 11 invention are short molecules containing a reactive cyclohexene group, which 12 may be dispersed in an appropriate medium for use. This may include the 13 use of short molecules (see also later) which can be dispersed within a 14 plastics resin or material. The ultimate result would be a plastics film or 15 material incorporating the oxygen scavenger. Of course, consideration would 16 need to be given to accessibility of the scavengers of the oxygen being 17 scavenged though this may rely on the porosity of the film (or film 18 layer/material) in which it is incorporated, or alternatively may be presented in 19 the manner of a coating with a reactive surface. 20 21 While the use of oxygen scavengers of varying sizes, (though typically those 22 of smaller size), dispersed through plastics materials is envisaged, oxygen 23 scavengers according to the present invention may also be used in other 24 ways. 25 26 For instance, they may be dispersed throughout non-plastics materials. This 27 may include inert and inorganic materials. This may also include other 28 liquids. It is envisaged that such embodiments of the present invention may 29 be used in applications such as sachets inserted into closed packages. The

use of oxygen scavengers in sachets and package inserts is documented in 1 2 the art and the same principles may be applied here. 3 4 Another means by which the present invention may be applied is through the 5 use of plastics resins incorporating the desired scavenging functionalities. 6 These resins, which for instance may include polyester resins, may be used 7 in the various manners by which resins are normally used. This may include 8 film production, resin coatings, as well as molding or extrusion techniques. 9 Another method by which the present invention may be implemented is the 10 formation or modification of polymers to contain the desired scavenging 11 functionalities and groups. In such cases, the film or plastics material itself 12 will possess oxygen scavenging properties. It is envisaged that such 13 materials may exist as layers in multi-layer films. Such polymers may also be 14 introduced as copolymers or blends in film and plastics manufacturing 15 methods. 16 17 The above instances of how embodiments of the present invention may be 18 used are illustrative only. It is noted that the use of oxygen scavenging 19 materials is known in the art, and that art may be drawn upon to further 20 expand the illustrative examples given within this specification. 21 22 Embodiments of the present invention based on cyclohexene groups appear 23 to afford significant advantage over the prior art. This advantage is in the 24 number and nature of the oxidation product once the scavenging is 25 completed. In the prior art, heavily reliance is made on straight chain alkenes, such as for instance fatty acids. The problem however, is that film 26 27 containing unsaturated compounds such as squalene or vegetable oils 28 produce large amounts of volatile aldehydes and ketones upon oxidation. 29 These tend to be released, or leach, from the plastics material over time,

usually find their way into the head space of the packaged material. The 1 2 presence of these foreign substances can represent a significant problem. 3 which the use of cyclohexene scavenging groups addresses at least partially. 4 5 In comparison, there are significantly less scission products from oxidation 6 reactions involving cyclohexene groups—the oxidation of the cyclohexene 7 group does not normally involve ring breakage. If the remainder of the molecule to which the cyclohexene group is attached is bound or linked to the 8 9 polymeric structure of the material in which it is incorporated or affixed, or 10 otherwise bound or held in place to the material through which it is dispersed 11 or incorporated, then there is little chance of there being any free scission 12 products able to find their way from the film or material structure. 13 14 Other aspects of the present invention to some extent parallel the prior art. 15 For instance, it is desirable that the oxygen scavenging materials are 16 relatively stable (with respect to scavenging) until required. In many cases, 17 catalysis and/or triggering of any reaction is required. Photo-initiators such as 18 benzophenone may be included. Initiating or triggering by electromagnetic 19 irradiation (often in the visible through UV regions) is convenient form of 20 triggering and already used in some types of film manufacture. It is also used 21 for triggering many prior art oxygen scavengers and thus employing these 22 features and techniques of the prior art with the present invention is 23 envisaged. 24 25 As for most other oxygen scavengers relying on alkenes, some form of 26 catalyst is also required for the oxygen scavenging reactions to proceed 27 effectively. Typically transition metal catalysts are used, including their salts, 28 complexes, and other compounds. These are well documented in the prior 29 art and may also be used with the present invention as appropriate. 30

1	
2	
3	According to another aspect of the present invention, there is provided an
4	oxygen scavenging polymer, substantially as described above, in which the
5	reactive extrusion process comprises a transesterification process.
6	•
7	
8	
9	According to one aspect of the present invention, there is provided an oxyger
10	scavenging composition for use in or with plastics materials, said scavenger
11	comprising or including at least one cyclohexene functionality as described
12	above.
13	· ·
14	According to another aspect of the present invention, there is provided an
15	oxygen scavenging composition, as described above, which produces only
16	low levels of volatile or extractable (from a plastics material in which it is
17	incorporated) products as a consequence of oxygen scavenging.
18	
19	According to another aspect of the present invention, there is provided an
20	oxygen scavenging composition, substantially as described herein which is
21	substantially stable with respect to reaction with oxygen until triggered by an
22	external event.
23	
24	According to another aspect of the present invention, there is provided an
25	oxygen scavenging composition, substantially as described above, wherein
26	the external event is irradiation by actinic radiation or electron beam radiation
27	
28	According to a further aspect of the present invention, there is provided an
29	oxygen scavenging composition including an oxygen scavenger, substantially

1,	as described above, which includes one or more trigger enhancing
2	components making the scavenger susceptible to triggering from an external
3	event.
4	
5	According to another aspect of the present invention, there is provided an
6	oxygen scavenging composition, substantially as described above, wherein a
7	trigger enhancing component is a photo initiator such as benzophenone or
8	substituted derivatives thereof.
9	
10	According to another aspect of the present invention, there is provided an
11	oxygen scavenging composition, substantially as described above, which
12	includes the presence of pone or more catalysts for the scavenging process.
13	
14	According to another aspect of the present invention, there is provided an
15	oxygen scavenging composition, substantially as described above, in which
16	the catalyst is a transition metal salt, compound or complex.
17	
18	According to another aspect of the present invention, there is provided an
19	oxygen scavenging composition, substantially as described above, which is in
20	the form of a plastic resin.
21	
22	According to another aspect of the present invention, there is provided an
23	oxygen scavenging composition, substantially as described above, in which
24	the plastic resin is a suitable for use in the manufacture of plastics films.
25	
26	According to another aspect of the present invention, there is provided an
27	oxygen scavenging composition, substantially as described above, in which
28	the plastic resin is a polyester resin.
29	

1 According to another aspect of the present invention, there is provided an 2 oxygen scavenging article comprising the oxygen scavenging composition 3 described above, where the scavenging component is present as a plastic 4 film or layer thereof. 5 6 According to another aspect of the present invention, there is provided an 7 oxygen scavenger or oxygen scavenging composition, substantially as 8 described above, when used as a polymeric material of a plastic film a layer 9 thereof, and/or a coating thereof, or in a plastic material. 10 11 According to another aspect of the present invention, there is provided an 12 oxygen scavenger or oxygen scavenging composition, substantially as 13 described above, when dispersed through a plastics film, a layer thereof, 14 and/or a coating thereon, or in a plastics material. 15 16 According to a further aspect of the present invention, there is provided an 17 oxygen scavenger or oxygen scavenging composition prepared from the 18 reaction of a tetrahydrophthalic anhydride or tetrahydrophthalic acid with at 19 least one of a diol, a hydroxy compound or polyhydroxy compound, in the 20 presence of or absence of an esterification catalyst. 21 22 According to a further aspect of the present invention, there is provided an 23 oxygen scavenger or oxygen scavenging composition prepared from the 24 reaction of a tetrahydrophthalic anhydride or tetrahydrophthalic acid with at 25 least one of a diol, a hydroxy compound or polyhydroxy compound, in the 26 presence of or absence of an esterification catalyst. 27 28 According to a further aspect of the present invention there is provided an 29 oxygen scavenger or oxygen scavenging composition prepared from an ester

1 or diester of a tetrahydrophthalic anhydride, in the presence of or absence of 2 a transesterification or esterification catalyst. 3 4 According to a further aspect of the present invention, there is provided an 5 oxygen scavenger or oxygen scavenging composition substantially as 6 described above, in which the anhydride comprises 1,2,3,6 tetrahydrophthalic 7 anhydride or tetrahydrophthalic anhydride monomers derived from butadiene. 8 2,3-Dimethyl-1,3-butadiene or isoprene. 9 10 According to a further aspect of the present invention, there is provided an 11 oxygen scavenging polymer including at least one cyclohexene group 12 prepared by a reactive extrusion process. 13 14 According to the present invention, a composition is provided comprising a 15 polymeric backbone, cyclic olefinic pendent groups and linking groups linking 16 the olefinic pendent groups to the polymeric backbone. 17 Also according to the present invention, an oxygen scavenging composition is 18 provided comprising a polymeric backbone, cyclic olefinic pendent groups, 19 linking groups linking the olefinic pendent groups to the polymeric backbone 20 and a transition metal catalyst. Also according to the present invention, an article of manufacture is provided 21 22 which is suitable as a container, the container inhibiting oxidation of contents 23 of the container by removing oxygen from the container and by inhibiting 24 ingress of oxygen into the container from outside the container, the article 25 comprising an oxygen scavenging composition which comprises a polymeric 26 backbone, cyclic olefinic pendent groups, linking groups linking the olefinic 27 pendent groups to the backbone, and a transition metal catalyst.

- 1 Also according to the present invention, a layer suitable for scavenging
- 2 oxygen is provided which comprises (a) a polymer backbone; (b) cyclic
- 3 olefinic pendent groups; (c) linking groups linking the backbone with the
- 4 pendent groups; and (d) a transition metal catalyst.
- 5 Also according to the present invention, a process of making a polymer
- 6 material is provided, the process being selected from the group consisting of
- 7 esterification, transesterification, amidation, transamidation and direct
- 8 polymerization, in which the oxygen scavenging packaging material
- 9 comprises a polymer backbone, cyclic olefinic pendent groups, linking groups
- 10 linking the backbone with the pendent groups.
- In a preferred embodiment of the invention, the polymeric backbone of the
- 12 above compositions, article, layer and process is ethylenic and the linking
- 13 groups are selected from the group consisting of:
- 14 -O-(CHR)_n-; -(C=O)-O-(CHR)_n-; -NH-(CHR)_n-; -O-(C=O)-(CHR)_n-;
- 15 -(C=O)-NH-(-CHR)₀-; and -(C=O)-O-CHOH-CH₂-O-;
- wherein R is hydrogen or an alkyl group selected from the group consisting of
- methyl, ethyl, propyl and butyl groups and where n is an integer in the range
- 18 from 1 to 12.
- 19 In a more preferred embodiment of the invention, the cyclic olefinic pendent
- 20 groups of the above compositions, article, layer and process have the
- 21 structure (II):

1

- where q_1 , q_2 , q_3 , q_4 , and r are selected from the group consisting of -H, -CH₃,
- 3 and $-C_2H_5$; and where m is $-(CH_2)_n$ with n being an integer in the range from 0
- 4 to 4; and wherein, when r is -H, at least one of q_1 , q_2 , q_3 and q_4 is -H.
- 5 In another preferred embodiment of the invention, the polymeric backbone of
- 6 the above compositions, article, layer and process comprises monomers
- 7 selected from the group consisting of ethylene and styrene.
- 8 In yet another preferred embodiment of invention, the cyclic olefinic pendent
- 9 groups of the above compositions, article, layer and process are grafted onto
- 10 the linking groups of the polymeric backbone by a esterification,
- 11 transesterification, amidation or transamidation reaction.
- 12 In still another preferred embodiment of the invention, the esterification,
- transesterification, amidation or transamidation reaction of the above
- 14 compositions, article, layer and process is a solution reaction or a reactive
- 15 extrusion.
- 16 In another preferred embodiment of the invention, the esterification.
- 17 transesterification, amidation or transamidation reaction of the above
- 18 compositions, article, layer and process is catalyzed by a catalyst selected
- 19 from the group consisting of strong non-oxidizing acids, tertiary amines,
- 20 Group I alkoxides, Group IVB alkoxides, and Group IVA organometallics.
- 21 In yet another preferred embodiment of invention, the catalyst of the above
- 22 compositions, article, layer and process is selected from a group consisting of
- 23 toluene sulfonic acid, sodium methoxide, tetrabutyl titanate, tetraisopropyl
- 24 titanate, tetra-n-propyl-titanate, tetraethyl titanate, 2-hydroxy-pyridine and
- 25 dibutyltin dilaurate.

26

- 1 In still another preferred embodiment of the invention, the polymeric
- 2 backbone, linking groups and cyclic olefin pendent groups of the above
- 3 compositions, article, layer and process comprise repeating units, each unit
- 4 having a structure (III) as follows:
- 5 (III)

- wherein P+T+ Q is 100 mol % of the total composition; P is greater than 0
- 11 mol % of the tot
- 12 al composition; Z is selected from the group consisting of an aryl group;
- 13 -(C=O)OR₁; -O(C=O)R₁; and an alkyl aryl group, structure (IV):

- where R₄ is selected from the group consisting of –CH₃, -C₂H₅, and -H; R₁ is
- selected from the group consisting of -H, -CH₃, -C₂H₅, -C₃H₇ and -C₄H₉; R₂ and
- 18 R₃ are selected from the group consisting of -H and -CH₃; X is selected from
- 19 the group consisting of -O-, -NH-, -(C=O)O-, -(C=O)NH-, -(C=O)S-, -O(C=O)-
- and -(CHR),-; ℓ is an integer in the range from 1 to 6; Y is -(CHR),-, where n is
- 21 an integer in the range from 0 to 12, R being selected from the group
- 22 consisting of –H, -CH₃ and -C₂H₅; where q₁, q₂, q₃, q₄, and r are selected from
- 23 the group consisting of -H, -CH₃, and -C₂H₅; and where m is -(CH₂)_n- and

- 1 where n is an integer in the range from 0 to 4; and wherein when r is -H, at
- 2 least one of q_1 , q_2 , q_3 and q_4 is -H.
- 3 In another preferred embodiment of the invention, the cyclic olefinic pendent
- 4 groups of the above compositions, article, layer and process are selected
- 5 from the group consisting of cyclohexene-4-methylene radical, 1-methyl
- 6 cyclohexene-4-methylene radical, 2-methyl cyclohexene-4-methylene radical,
- 7 5-methyl cyclohexene-4-methylene radical, 1,2-dimethyl cyclohexene-4-
- 8 methylene radical, 1,5-dimethyl cyclohexene-4-methylene radical,
- 9 2,5-dimethyl cyclohexene-4-methylene radical, 1,2,5-trimethyl cyclohexene-4-
- 10 methylene radical, cyclohexene-4-ethylene radical, 1-methyl cyclohexene-4-
- 11 ethylene radical, 2-methyl cyclohexene-4-ethylene radical, 5-methyl
- 12 cyclohexene-4-ethylene radical, 1,2-dimethyl cyclohexene-4-ethylene radical.
- 13 1,5-dimethyl cyclohexene-4-ethylene radical, 2,5-dimethyl cyclohexene-4-
- 14 ethylene radical, 1,2,5-trimethyl cyclohexene-4-ethylene radical, cyclohexene-
- 15 4-propylene radical, 1-methyl cyclohexene-4-propylene radical, 2-methyl
- 16 cyclohexene-4-propylene radical, 5-methyl cyclohexene-4-propylene radical,
- 17 1,2-dimethyl cyclohexene-4-propylene radical, 1,5-dimethyl cyclohexene-4-
- 18 propylene radical, 2,5-dimethyl cyclohexene-4-propylene radical,
- 19 1,2,5-trimethyl cyclohexene-4-propylene radical, cyclopentene-4-methylene
- 20 radical, 1-methyl cyclopentene-4-methylene radical, 3-methyl cyclopentene-4-
- 21 methylene radical, 1,2-dimethyl cyclopentene-4-methylene radical,
- 22 3,5-dimethyl cyclopentene-4-methylene radical, 1,3-dimethyl cyclopentene-4-
- 23 methylene radical, 2,3-dimethyl cyclopentene-4-methylene radical,
- 24 1,2,3-trimethyl cyclopentene-4-methylene radical, 1,2,3,5-tetramethyl
- 25 cyclopentene-4-methylene radical, cyclopentene-4-ethylene radical, 1-methyl
- 26 cyclopentene-4-ethylene radical, 3-methyl cyclopentene-4-ethylene radical,
- 27 1,2-dimethyl cyclopentene-4-ethylene radical, 3,5-dimethyl cyclopentene-4-
- 28 ethylene radical, 1,3-dimethyl cyclopentene-4-ethylene radical, 2,3-dimethyl
- 29 cyclopentene-4-ethylene radical, 1,2,3-trimethyl cyclopentene-4-ethylene

1 radical, 1,2,3,5-tetramethyl cyclopentene-4-ethylene radical, cyclopentene-4-2 propylene radical, 1-methyl cyclopentene-4-propylene radical, 3-methyl 3 cyclopentene-4-propylene radical, 1,2-dimethyl cyclopentene-4-propylene 4 radical, 3,5-dimethyl cyclopentene-4-propylene radical, 1,3-dimethyl 5 cyclopentene-4-propylene radical, 2,3-dimethyl cyclopentene-4-propylene 6 radical, 1,2,3-trimethyl cyclopentene-4-propylene radical, and 7 1,2,3,5-tetramethyl cyclopentene-4-propylene radical. 8 In yet another preferred embodiment of the invention, the composition of the 9 above compositions, article, layer and process is a ethylene/methyl 10 acrylate/cyclohexenyl methyl acrylate terpolymer, a cyclohexenyl methyl 11 acrylate/ethylene copolymer, a cyclohexenyl methyl methacrylate/styrene 12 copolymer, a cyclohexenyl methyl acrylate homopolymer or a methyl 13 acrylate/cyclohexenyl methyl acrylate copolymer. 14 15 In another preferred embodiment of the invention, the odor and taste 16 characteristics of products packaged with material comprised of the above 17 compositions, article, layer and process are not adulterated as a result of 18 oxidation of the composition. 19 20 In still another preferred embodiment of the invention, there is no significant 21 fragmentation of the olefinic pendent groups and linking groups from the 22 polymeric backbone as a result of oxidation of the above compositions, 23 article, layer and process. 24 25 In yet another preferred embodiment of the invention, the transition metal 26 catalyst of the above oxygen scavenging composition, article of manufacture, 27 layer and process is a metal salt.

1	In still another preferred embodiment of the invention, the metal in the metal
2	salt of the above oxygen scavenging composition, article of manufacture,
3	layer and process is cobalt.
4	In still another preferred embodiment of the invention, the metal salt of the
5	above oxygen scavenging composition, article of manufacture, layer and
6	process is selected from the group consisting of cobalt neodecanoate, cobalt
7	2-ethylhexanoate, cobalt oleate and cobalt stearate.
8	
9	In yet another preferred embodiment of the invention, the composition of the
10	above oxygen scavenging composition, article of manufacture, layer and
11	process further comprises at least one triggering material to enhance initiation
12	of oxygen scavenging.
13	
14	In still another preferred embodiment of the invention, the triggering material
15	of the above oxygen scavenging composition, article of manufacture, layer
16	and process is a photo initiator.
17	
18	In a preferred embodiment of the invention, the above article of manufacture
19	is a package.
20	·
21	In another preferred embodiment of invention, the package article of the
22	above article of manufacture comprises a flexible film having a thickness of at
23	most 10 mil or a flexible sheet having a thickness of at least 10 mil.
24	
25	In yet another preferred embodiment of the invention, the oxygen scavenging
26	system of the package article of the above article of manufacture comprises
27	at least one additional layer selected from among oxygen barrier layers,
28	polymeric selective layers, and heat seal layers.
29	

In still another preferred embodiment of the invention, the above article of 1 manufacture is a package with a food product located within the package. 2 3 In yet another preferred embodiment of the invention, the above article of 4 manufacture is a package for packaging a cosmetic, chemical, electronic 5 device, pesticide or a pharmaceutical composition. 6 7 In still another preferred embodiment of the invention, a multi-layer film 8 comprises the article of the above article of manufacture and the film has at 9 least one additional functional layer. 10 11 In yet another preferred embodiment of the invention, the multi-layer film of 12 the above article of manufacture has at least one additional layer selected 13 from among oxygen barrier layers, polymeric selective barrier layers, 14 structural layers and heat seal layers. 15 16 In still another preferred embodiment of the invention, the multi-layer film of 17 the above article of manufacture has at least one additional layer which is an oxygen barrier layer. 18 19 20 In yet another preferred embodiment of the invention, the multi-layer film of 21 the above article of manufacture further comprises at least one polymeric 22 selective barrier layer. 23 24 In still another preferred embodiment of the invention, the multi-layer film of 25 the above article of manufacture further comprises at least one heat seal 26 layer. 27 28 In yet another preferred embodiment of the invention, the multi-layer film of 29 the above article of manufacture further comprises at least one structural 30 layer.

1	m sun another presented embodiment of the invention, the above afticle of
2	manufacture is a rigid container, sealing gasket, patch, container closure
3	device, bottle cap, bottle cap insert or molded or thermoformed shape.
4	
5	In yet another preferred embodiment of the invention, the molded or
6	thermoformed shape of the above article of manufacture is a bottle or tray.
7	
8	In still another preferred embodiment of the invention, the above layer in
9	addition comprises polymeric diluent.
10	
11	In yet another preferred embodiment of the invention, the diluent of the above
12	layer is a thermoplastic polymer.
13	·
14	In still another preferred embodiment of the invention, the above layer is
15	adjacent to one or more additional layers.
16	
17	In still another preferred embodiment of the invention, at least one of the
18	additional layers adjacent to the above layer is an oxygen barrier.
19	
20	In still another preferred embodiment of the invention, the oxygen barrier of
21	the above layer comprises a member of the group consisting of poly(ethylene-
22	vinyl alcohol), polyacrylonitrile, poly(vinyl chloride), polyamides,
23	poly(vinylidene dichloride), poly(ethylene terephthalate), silica, metal foil and
24	metalized polymeric films.
25	
26	In still another preferred embodiment of the invention, the one or more of said
27	additional layer or layers of the above layer is coextruded with the above
28	layer.
29	

1	In y	et another preferred embodiment of the invention, the one or more of said
2	add	itional layer or layers of the above layer is laminated onto the above layer
3		
4	In s	till another preferred embodiment of the invention, the one or more of said
5	add	itional layer or layers of the above layer is coated onto the above layer.
6		
7	In y	et another preferred embodiment of the invention, the above layer is
8	flex	ible.
9		
10	In s	till another preferred embodiment of the invention, the above layer is
11	tran	sparent.
12		
13	In y	et another preferred embodiment of the invention, an article for packaging
14	whe	erein the article comprises the above layer.
15		
16	In y	et another preferred embodiment of the invention, the above process of
17	mak	king the oxygen scavenging packaging material comprises the steps of:
18	(a)	selecting polymers from the group consisting of styrene/maleic
19		anhydride, ethylene/maleic anhydride, ethylene/acrylic acid,
20		ethylene/methacrylic acid, acrylic acid, methacrylic acid,
21		styrene/methacrylic acid, ethylene/methyl acrylate, ethylene/ethyl
22		acrylate, ethylene/butyl acrylate, methyl methacrylate, methyl acrylate,
23		and styrene/methyl methacrylate to form a mixture and combining the
24		polymers with an esterifying/transesterifying compound selected from
25		the group consisting of cyclohexene-4-methanol, 1-methyl cyclohexene-
26		4-methanol, 2-methyl cyclohexene-4-methanol, 5-methyl cyclohexene-4
27		methanol, 1,2-dimethyl cyclohexene-4-methanol, 1,5-dimethyl
28		cyclohexene-4-methanol, 2,5-dimethyl cyclohexene-4-methanol,

1		1,2,5-trimethyl cyclohexene-4-methanol, cyclohexene-4-ethanol,
2		1-methyl cyclohexene-4-ethanol, 2-methyl cyclohexene-4-ethanol,
3		5-methyl cyclohexene-4-ethanol, 1,2-dimethyl cyclohexene-4-ethanol,
4		1,5-dimethyl cyclohexene-4-ethanol, 2,5-dimethyl cyclohexene-4-
5		ethanol, 1,2,5-trimethyl cyclohexene-4-ethanol, cyclohexene-4-propanol
6		1-methyl cyclohexene-4-propanol, 2-methyl cyclohexene-4-propanol,
7		5-methyl cyclohexene-4-propanol, 1,2-dimethyl cyclohexene-4-propanol
8		1,5-dimethyl cyclohexene-4-propanol, 2,5-dimethyl cyclohexene-4-
9		propanol, 1,2,5-trimethyl cyclohexene-4-propanol, cyclopentene-4-
10		methanol, 1-methyl cyclopentene-4-methanol, 3-methyl cyclopentene-4-
11		methanol, 1,2-dimethyl cyclopentene-4-methanol, 3,5-dimethyl
12		cyclopentene-4-methanol, 1,3-dimethyl cyclopentene-4-methanol,
13		2,3-dimethyl cyclopentene-4-methanol, 1,2,3-trimethyl cyclopentene-4-
14		methanol, 1,2,3,5-tetramethyl cyclopentene-4-methanol, cyclopentene-
15		4-ethanol, 1-methyl cyclopentene-4-ethanol, 3-methyl cyclopentene-4-
16		ethanol, 1,2-dimethyl cyclopentene-4-ethanol, 3,5-dimethyl
17		cyclopentene-4-ethanol, 1,3-dimethyl cyclopentene-4-ethanol,
18		2,3-dimethyl cyclopentene-4-ethanol, 1,2,3-trimethyl cyclopentene-4-
19		ethanol, 1,2,3,5-tetramethyl cyclopentene-4-ethanol, cyclopentene-4-
20		propanol, 1-methyl cyclopentene-4-propanol, 3-methyl cyclopentene-4-
21		propanol, 1,2-dimethyl cyclopentene-4-propanol, 3,5-dimethyl
22		cyclopentene-4-propanol, 1,3-dimethyl cyclopentene-4-propanol,
23		2,3-dimethyl cyclopentene-4-propanol, 1,2,3-trimethyl cyclopentene-4-
24		propanol, and 1,2,3,5-tetramethyl cyclopentene-4-propanol;
25	(b)	heating the polymers and esterifying/transesterifying compounds
26		selected in (a) to form a polymer melt;
27	(c)	processing the melt in an extruder under esterification/transesterification
28		conditions with esterification/transesterification catalysts and

1	antioxidants protecting the melt from oxidation during extrusion, so that
2	the polymer melt undergoes esterification of polymeric anhydrides with
3	cyclic olefin pendent groups, esterification of polymeric acids with cyclic
4	olefin pendent groups or exchange of alkyl groups of polymeric esters
5	with cyclic olefin pendent groups; and

- 6 (d) removing volatile organic products and by-products from the melt.
- 7 In still another preferred embodiment of the invention, the above process of
- 8 making the oxygen scavenging packaging material comprises the steps of:
- 9 selecting polymers from the group consisting of styrene/maleic 10 anhydride, ethylene/maleic anhydride, ethylene/acrylic acid, 11 ethylene/methacrylic acid, acrylic acid, methacrylic acid, 12 styrene/methacrylic acid, ethylene/methyl acrylate, ethylene/ethyl acrylate, ethylene/butyl acrylate, methyl methacrylate, methyl acrylate, 13 14 and styrene/methyl methacrylate to form a mixture and combining the 15 polymers with an amidizing/transamidizing compound selected from the 16 group consisting of cyclohexene-4-methyl amine, 1-methyl cyclohexene-17 4-methyl amine, 2-methyl cyclohexene-4-methyl amine, 5-methyl 18 cyclohexene-4-methyl amine, 1,2-dimethyl cyclohexene-4-methyl amine, 19 1,5-dimethyl cyclohexene-4-methyl amine, 2,5-dimethyl cyclohexene-4-20 methyl amine, 1,2,5-trimethyl cyclohexene-4-methyl amine. 21 cyclohexene-4-ethyl amine, 1-methyl cyclohexene-4-ethyl amine, 22 2-methyl cyclohexene-4-ethyl amine, 5-methyl cyclohexene-4-ethyl 23 amine, 1,2-dimethyl cyclohexene-4-ethyl amine, 1,5-dimethyl 24 cyclohexene-4-ethyl amine, 2,5-dimethyl cyclohexene-4-ethyl amine, 25 1,2,5-trimethyl cyclohexene-4-ethyl amine, cyclohexene-4-propyl amine, 26 1-methyl cyclohexene-4-propyl amine, 2-methyl cyclohexene-4-propyl 27 amine, 5-methyl cyclohexene-4-propyl amine, 1,2-dimethyl cyclohexene-28 4-propyl amine, 1,5-dimethyl cyclohexene-4-propyl amine, 2,5-dimethyl

1		cyclohexene-4-propyl amine, 1,2,5-trimethyl cyclohexene-4-propyl
2		amine, cyclopentene-4-methyl amine, 1-methyl cyclopentene-4-methyl
3		amine, 3-methyl cyclopentene-4-methyl amine, 1,2-dimethyl
4		cyclopentene-4-methyl amine, 3,5-dimethyl cyclopentene-4-methyl
5		amine, 1,3-dimethyl cyclopentene-4-methyl amine, 2,3-dimethyl
6		cyclopentene-4-methyl amine, 1,2,3-trimethyl cyclopentene-4-methyl
7		amine, 1,2,3,5-tetramethyl cyclopentene-4-methyl amine, cyclopentene-
8		4-ethyl amine, 1-methyl cyclopentene-4-ethyl amine, 3-methyl
9		cyclopentene-4-ethyl amine, 1,2-dimethyl cyclopentene-4-ethyl amine,
10		3,5-dimethyl cyclopentene-4-ethyl amine, 1,3-dimethyl cyclopentene-4-
11		ethyl amine, 2,3-dimethyl cyclopentene-4-ethyl amine, 1,2,3-trimethyl
12		cyclopentene-4-ethyl amine, 1,2,3,5-tetramethyl cyclopentene-4-ethyl
13		amine, cyclopentene-4-propyl amine, 1-methyl cyclopentene-4-propyl
14		amine, 3-methyl cyclopentene-4-propyl amine, 1,2-dimethyl
15		cyclopentene-4-propyl amine, 3,5-dimethyl cyclopentene-4-propyl
16		amine, 1,3-dimethyl cyclopentene-4-propyl amine, 2,3-dimethyl
17		cyclopentene-4-propyl amine, 1,2,3-trimethyl cyclopentene-4-propyl
18		amine, and 1,2,3,5-tetramethyl cyclopentene-4-propyl amine;
19	(b)	heating the polymers and amidizing/transamidizing compounds selected
20		in (a) to form a polymer melt;
21	(c)	processing the melt in an extruder under amidation/transamidation
22	(0)	conditions with amidation/transamidation catalysts and antioxidants
23		•
24		protecting the melt from oxidation during extrusion, so that the polymer
		melt undergoes amidation of polymeric anhydrides with cyclic olefin
25		pendent groups, amidation of polymeric acids with cyclic olefin pendent
26		groups or exchange of alkyl groups of polymeric esters with cyclic olefin
27		pendent groups; and

(d) removing volatile organic products and by-products from the melt.

1 In yet another preferred embodiment of the invention, the above process of 2 making of the oxygen scavenging packaging material comprises the steps of: 3 adding to an autoclave, ethylene and a vinyl monomer comprising a 4 pendent cyclohexene; 5 6 stirring the ethylene and the vinyl monomer in the autoclave to achieve a 7 mixture; 8 9 (c) adding a polymerization initiator before, during or after the stirring step; 10 11 polymerizing the mixture to achieve a polymer; and 12 13 (e) isolating and purifying the polymer. 14 15 In still another embodiment of the invention, in the above process, in step (a), 16 an alpha-olefin is added to the autoclave along with the ethylene and the vinyl 17 monomer and, in step (b), the alpha-olefin is stirred with the ethylene and the 18 vinyl monomer to achieve the mixture. 19 20 The present invention relates to a non-odorous oxygen scavenging polymer 21 composition comprising: (1) monomers derived from cyclic hydrocarbon 22 moieties having at least one cyclic allylic or cyclic benzylic hydrogen and (2) a 23 transition metal oxidation catalyst. The present invention also relates to a 24 rigid container for food or beverage, the container being molded from a resin 25 comprising the above-described non-odorous oxygen scavenging polymer 26 composition. The present invention also relates to the above-described rigid 27 container further comprising a tinted ultraviolet protection layer, which may or 28 may not be the food contact layer, located between the layer comprising the 29 non-odorous oxygen scavenging composition and the inside of the rigid 30 container.

1 In a preferred embodiment of the above non-odorous oxygen scavenging 2 polymer composition, wherein the composition comprises a vinyl polymer 3 selected from the group consisting of ethylene polymer, ethylene copolymer. 4 propylene polymer, propylene copolymer, styrene polymer, styrene copolymer 5 and mixtures thereof. 6 . 7 In another preferred embodiment of the above non-odorous oxygen 8 scavenging polymer composition, the composition comprises condensation 9 polymers selected from the group consisting of polyesters, polyamides. 10 polycarbonate, polyurethane, polyureas and polyether. 11 12 In a more preferred embodiment of the above composition comprising 13 condensation polymers, the composition is thermoplastic. 14 15 In another more preferred embodiment of the above composition comprising 16 condensation polymers, the composition is thermoset. 17 18 In yet another more preferred embodiment of the above composition 19 comprising condensation polymers, the composition is a multilayered 20 structure with other layers being an aromatic polyester or copolyester 21 selected from the group consisting of polyethylene terephthalate, 22 polyethylene naphthalate, polypropylene terephthalate, polybutylene 23 terephthalate, polyethylene isophthalate, polycyclohexanedimethanol 24 terephthalate, polybutylene naphthalate, polycyclohexanedimethanol 25 naphthalate, and copolymers and blends thereof. 26 27 In still another more preferred embodiment of the above composition 28 comprising condensation polymers, the composition is a multilayered 29 structure with other layers being polyamides or copolyamides selected from 30 the group consisting of Nylon 6, Nylon 66, Nylon 610 and mixtures thereof.

1	In yet another more preferred embodiment of the above composition
2	comprising condensation polymers, the composition is a multilayered
3	structure with other layers being bisphenol A carbonate.
4	
5	In yet another more preferred embodiment of the above composition
6	comprising condensation polymers, the composition is a multilayered
7	structure with other layers being vinylic polymers or copolymers selected
8	from the group consisting of ethylene polymer, ethylene copolymer, propylene
9	polymer, propylene copolymer, styrene polymer, styrene copolymer, acrylate
10	polymer, acrylate copolymer, vinyl chloride polymer, vinyl chloride copolymer,
11	divinyl chloride polymer, divinyl chloride copolymer, fluorinated vinyl polymer,
12	fluorinated vinyl copolymer and mixtures thereof.
13	
14	In still another more preferred embodiment of the above composition
15	comprising condensation polymers, the composition is blended with an
16	aromatic polyester or copolyester selected from the group consisting of
17	polyethylene terephthalate, polyethylene naphthalate, polypropylene
18	terephthalate, polybutylene terephthalate, polyethylene isophthalate,
19	polycyclohexandedimethanol terephthalate, polybutylene naphthalate,
20	polycyclohexanedimethanol naphthalate, and copolymers and blends thereof.
21	
22	In yet another more preferred embodiment of the above composition-
23	comprising condensation polymers, the composition is blended with
24	polyamides or copolyamides selected from the group consisting of Nylon 6,
25	Nylon 66, Nylon 610 and mixtures thereof.
26	
27	In still another more preferred embodiment of the above composition
28	comprising condensation polymers, the composition is blended with bisphenol
29	A polycarbonate.

1 In yet another more preferred embodiment of the above composition

2 comprising condensation polymers, the composition being a blend comprising

3 vinylic polymers or copolymers selected from the group consisting of ethylene

4 polymer, ethylene copolymer, propylene polymer, propylene copolymer,

5 styrene polymer, styrene copolymer, acrylate polymer, acrylate copolymer,

6 vinyl chloride polymer, vinyl chloride copolymer, divinyl chloride polymer.

7 divinyl chloride copolymer, fluorinated vinyl polymer, fluorinated vinyl

8 copolymer and mixtures thereof.

9

11

12

13

14

15

In a more preferred embodiment of the above composition comprising condensation polymers, the composition is laminated or adhering onto a substrate selected from the group consisting of paper, foil, high temperature film, metallized film, polyamide films, ethylene vinyl alcohol film, silica coated film, nylon/EVOH/nylon, oriented polypropylene, polyester film, polyethylene, polypropylene, polyester, oriented polyethylene terephthalate and cellophane.

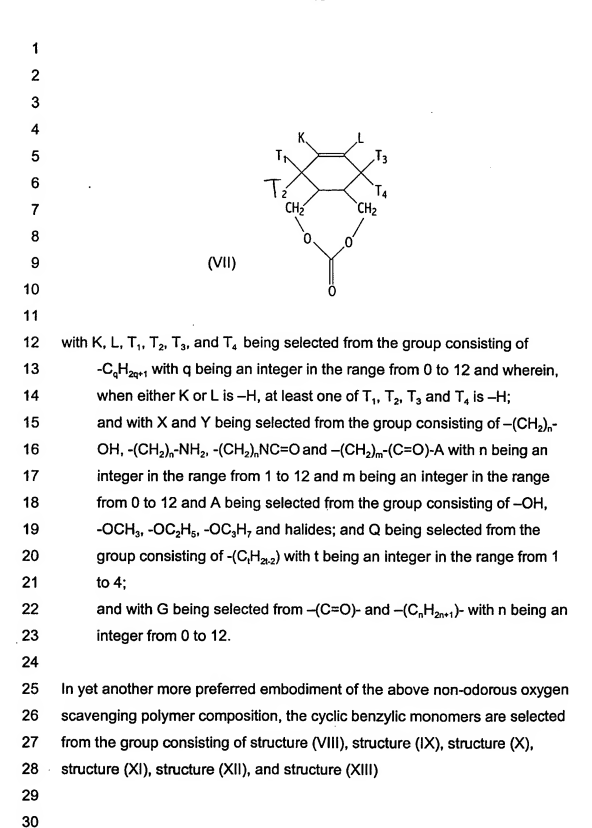
16 17

In another preferred embodiment of the above non-odorous oxygen scavenging polymer composition, the cyclic allylic monomers are selected from the group consisting of structure (VI), structure (VI) and structure (VII):

19 20

18

21
22
23
24
25
26
27
28
29
30
$$(V)$$



where X and Y are selected from the group consisting of $-(CH_2)_n$ -OH, $-(CH_2)_n$ -NH₂ and $-(CH_2)_m$ -(C=O)-R₁ with n being an integer in the range

1	from 1 to 12, and with m being an integer in the range from 0 to 12 and
2	with R₁ being selected from the group consisting of –OH, -OCH₃,
3	-OC₂H₅, -OC₃H₁ and halides;
4	with T_1 , T_2 , T_3 , and T_4 being selected from the group consisting of
5	$-C_qH_{2q+1}$ with q being an integer in the range from 0 to 12 and at least
6	one of T_1 , T_2 , T_3 and T_4 being $-H$;
7	and with X and Y being selected from the group consisting of -
8	$(CH_2)_n$ -OH, - $(CH_2)_n$ -NH ₂ , - $(CH_2)_n$ NC=O, and - $(CH_2)_m$ - $(C=O)$ -A with n
9	being an integer in the range from 1 to 12, and m being an integer in the
10	range from 0 to 12 and A being selected from the group consisting of
11	-OH, -OCH ₃ , -OC ₂ H ₅ , -OC ₃ H ₇ and halides; and Z being selected from the
12	group consisting of -(C_tH_{2t-2})-, -O-, -NR ₂ -, -S-, with t being an integer in
13	the range from 1 to 4 and R ₂ being selected from the group consisting of
14	–OH, -OCH₃, -OC₂H₅, -OC₃H₁ and halides;
15	and with G being selected from –(C=O)- and -(C_nH_{2n+1})- with n being an
16	integer from 0 to 12.
17	In still another more preferred embodiment, the composition of the resin of the
18	above-described rigid container is a single layer.
19	
20	In yet another more preferred embodiment, the composition of the resin of the
21	above-described rigid container is multilayered.
22	
23	In yet another more preferred embodiment, the composition of the resin of the
24	above-described rigid container comprises an outer air contact layer and an
25	inner oxygen scavenging layer.
26	
27	In still another more preferred embodiment, the outer air contact layer of the
28	composition of the resin of the above-described rigid container comprises an
29	oxygen barrier resin selected from the group consisting of polyethylene

WO 99/48963

1 terephthalate, polyethylene naphthalate and a mixture of polyethylene 2 terephthalate and polyethylene naphthalate. 3 4 In yet another more preferred embodiment, the composition of the resin of the 5 above-described rigid container further comprises at least one of an inner 6 food contact layer, a tie layer, and a tinted ultraviolet protection layer. 7 8 In still another more preferred embodiment, the inner food contact layer of the 9 composition of the resin of the above-described rigid container comprises an 10 oxygen barrier resin selected from the group consisting of polyethylene 11 terephthalate, polyethylene naphthalate and a mixture of polyethylene 12 terephthalate and polyethylene naphthalate. 13 14 In yet another more preferred embodiment, the oxygen scavenging of the 15 composition of the resin of the above-described rigid container is initiated by 16 moisture or actinic radiation. 17 In still another more preferred embodiment, the transition metal catalyst of the 18 composition of the resin of the above-described rigid container is a metal salt. 19 20 In yet another more preferred embodiment, the metal in the metal salt of the 21 transition metal catalyst of the composition of the resin of the above-22 described rigid container is cobalt. 23 24 In still another more preferred embodiment, the metal salt of the transition 25 metal catalyst of the composition of the resin of the above-described rigid 26 container is selected from the group consisting of cobalt neodecanoate, 27 cobalt 2-ethylhexanoate, cobalt oleate and cobalt stearate. 28

1	in yet another more preferred embodiment, the composition of the resin of the
2	above-described rigid container further comprises at least one triggering
3	material to enhance initiation of oxygen scavenging.
4	
5	In still another more preferred embodiment, the triggering material of the resin
6	of the composition of the above-described rigid container is a photoinitiator.
7	
8	In yet another more preferred embodiment, the photoinitiator of the resin of
9	the composition of the above-described rigid container has an ultraviolet
10	absorption window above 320 nm.
11	
12	In still another more preferred embodiment, the above-described rigid
13	container is suitable for packaging oxygen sensitive drinks for extended
14	freshness and shelf life.
15	
16	In yet another more preferred embodiment, the above-described rigid
17	container is suitable for packaging beer.
18	·
19	DESCRIPTION OF THE DRAWINGS
20	Figure 1 is a schematic showing the overall process leading to the
21	transesterification of ethylene methyl acrylate copolymers (EMAC) to give
22	modified EMAC having cyclic pendent olefins.
23	Figure 2 is a graph comparatively plotting percent oxygen in headspace at
24	·
25	4°C (initially at 1% oxygen) against time in days for two 3-layer film extrusions
26	based on Dowlex® 3010/EMCM/Dowlex® 3010 films (EMCM being an
	acronym for ethylene/methyl acrylate/cyclohexenyl methyl acrylate terpolymer
27	also referred to as poly(ethylene/methyl acrylate/cyclohexene-methyl
28	acrylate)), both including the EMCM inner layer and one of them having

1	50 ppm of a non-volatile antioxidant Irganox® 1010 in the EMCM layer and
2	one of them having 100 ppm Irganox® 1010 in the EMCM layer.
3	Figure 3 is a graph comparatively plotting percent oxygen in headspace at
4	4°C (initially at 1% oxygen) against time in days for an EMCM film and two
5	.EBAC blended EMCM films, one of them having 3:1 EBAC:EMCM and one of
6	them having 1:1 EBAC:EMCM.
7	Figure 4 is a graph comparatively plotting the oxygen scavenging rates and
8	capacities at 25°C in which the initial headspace oxygen was 21% (air) for an
9	EMCM film and a 2:1 EBAC:EMCM film.
10	Figure 5 is a graph showing the taste ratings in a comparative taste test
11	between food stored in two oxygen scavenging packages (EMCM and SBS)
12	and a control package (no oxygen scavenger).
13	
14	
15	
16	DETAILED DESCRIPTION OF THE INVENTION
17	
18	We have found that materials containing certain cyclohexenyl functionalities
19	are excellent oxygen absorbers when compounded with a transition metal sal
20	and optionally a photoinitiator, and that when these materials oxidize they
21	produce very low levels of oxidation byproducts. This is in marked contrast to
22	the known art, where excellent oxygen absorbers can be obtained from the
23	use of linear unsaturated compounds compound with a transition metal slat,
24	and a photoinitiator, but where the levels of oxidation byproducts are
25	excessively high. It is thought that this improvement is obtained because mile
26	oxidation of cyclohexene does not break bonds on the ring structure whilst
27	oxidation of linear unsaturated material such as linoleic acid or vegetable oil

28

(b)

a transition metal catalyst;

1 under similar conditions produces smaller molecules by chain scission. When 2 incorporated into polymers, the cyclohexene containing systems are found to 3 produce considerably less volatile byproducts than the linear unsaturated 4 materials. 5 The compositions of this invention are significantly cleaner than those 6 described in the prior art, they do not require the use of high levels of adjuncts 7 to absorb the undesirable byproducts. Such absorbent additives are known in 8 the art, for example see U.S. 5,834,079 and U.S. 08/857,276. It is also well 9 known in the art that such additives (zeolites and silicas) adversely affect the 10 haze and clarity of packaging structures. 11 12 The oxygen scavenging compositions consist of: 13 a polymer or lower molecular weight material containing substituted (a) 14 cyclohexene functionality according to the following structure (I): 15 16 17 18 **(l)** 19 20 21 22 23 24 where A may be hydrogen or methyl and either one or two of the B groups is a heteroatom containing linkage which attaches the cyclohexene ring to 25 the said material. The remaining B groups are hydrogen or methyl; 26

(c) an optional photoinitiator.

3 4

5

6

7 8

9

10

11

12

The compositions may be polymeric in nature or they may be lower molecular weight materials. In either case, they may be blended with further polymers or other additives. In the case of low molecular weight materials they will most likely be compounded with a carrier resin before use. The following examples represent some applications of various embodiments of the present invention currently envisaged by the patentee. These examples are not meant to be limiting nor exhaustive but merely illustrative of how the present invention may be used, or applied to address problems associated with the prior art.

13

14 15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

The compositions of this invention can be used in a wide range of packaging materials, and are not restricted to flexible packaging films and articles such as pouches produced from such films. The compositions may also be used in the preparation of rigid and semi rigid packaging materials. Typical rigid and semi rigid articles include plastic, paper or cardboard cartons, bottles such as juice containers, thermoformed trays, or cups with wall thicknesses of about 100 to 2000 microns. The walls of such articles comprise single or multiple layers of materials. The compositions can be used as the sole polymeric material from which one or more layers of a film are formed (i.e., the film can be a multilayer film having, for example, a gas barrier layer, a seal layer, etc.), it can be blended with other polymeric oxygen scavenging agents (such as polybutadiene) or it can be blended with one or more diluent polymers which are known to be useful in the formation of packaging film materials and which often can render the resultant film more flexible and/or processable. Suitable diluent polymers include, but are not limited to, polyethylene such as, for example, low density polyethylene, very low density polyethylene, ultra-low

1	density polyethylene, high density polyethylene, and linear low density
2	polyethylene; pólyesters such as, for example, polyethylene terephthalate
3	(PET); polyvinyl chloride (PVC); polyvinylidene chloride (PVDC); and ethylene
4	copolymers such as ethylene/vinyl acetate copolymer, ethylene/alkyl
5	(meth)acrylate copolymers, ethylene/(meth)acrylic acid copolymers, and
6	ionomers. Blends of different diluent polymers also can be used.
7	
8	The compositions of this invention can also be used in non integral packaging
9	components such as coatings, bottle cap liners, adhesive and non adhesive
10	sheet inserts, coupons, gaskets, sealants or fibrous mass inserts.
11	
12	Generally, the foregoing diluent polymers are semi-crystalline materials.
13	Advantageously, the polymeric component of the composition of the present
14	invention can be crystalline or semi-crystalline at ambient conditions and,
15	accordingly, can be especially compatible with such diluent polymers.
16	Selection of a particular diluent polymer(s) depends largely on the article to
17	be manufactured and the end use thereof. For instance, certain polymers are
18	known by the ordinarily skilled artisan to provide clarity, cleanliness, barrier
19	properties, mechanical properties, and/or texture to the resultant article.
2Ó	
21	In combination with the polymeric component, the oxygen scavenging
22	composition of the present invention includes a transition metal compound as
23	an oxygen scavenger catalyst. The transition metal catalyst can be a slat
24	which includes a metal selected from the first, second, or third transition
25	series of the Periodic Table. The metal preferably is Rh, Ru, or one of the
26	elements in the series of Sc to Zn (i.e., Sc, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, and
27	Zn), more preferably at least one of Mn, Fe, Co, Ni, and Cu, and most
28	preferably Co. Suitable anions for such salts include, but are not limited to,
29	chloride, acetate, oleate, stearate, palmitate, 2-ethylhexanoate,
30	neodecanoate, and naphthenate. Representative salts include cobalt (II)

1 2-ethylhexanoate, cobalt oleate, and cobalt (II) neodecanoate. (The metal 2 salt also can be an ionomer, in which case a polymeric counterion is 3 employed.) 4 5 When used in forming a packaging article, the oxygen scavenging 6 composition of the present invention can include only the above-described 7 polymers and a transition metal catalyst. However, photoinitiators can be 8 added to further facilitate and control the initiation of oxygen scavenging 9 properties. Adding a photoinitiator or a blend of photoinitiators to the oxygen 10 scavenging composition can be preferred, especially where antioxidants have 11 been added to prevent premature oxidation of the composition during 12 processing and storage. 13 14 Suitable photoinitiators are known to those skilled in the art. See, e.g., PCT 15 Publication WO 97/07161, WO 97/44364, WO 98/51758, and WO 98/51759. 16 the teachings of which are incorporated herein by reference as if set forth in 17 full. Specific examples of suitable photoinitiators include, but are not limited 18 to, benzophenone, and its derivatives, such as methoxybenzophenone, 19 dimethoxybenzophenone, dimethylbenzophenone, diphenoxybenzophenone, 20 allyloxybenzophenone, diallyloxybenzophenone, dodecyloxybenzophenone, 21 dibenzosuberone, 4,4'-bis(4-isopropylphenoxy)benzophenone, 22 4-morpholinobenzophenone, 4-aminobenzophenone, tribenzoyl 23 triphenylbenzene, tritoluoyl triphenylbenzene, 4,4'-bis(dimethylamino)-24 benzophenone, acetophenone and its derivatives, such as, o-methoxy-25 acetophenone, 4'-methoxyacetophenone, valerophenone, hexanophenone, 26 α-phenyl-butyrophenone, p-morpholinopropiophenone, benzoin and its 27 derivatives, such as, benzoin methyl ether, benzoin butyl ether, benzoin 28 tetrahydropyranyl ether, 4-o-morpholinodeoxybenzoin, substituted and 29 unsubstituted anthraquinones, α -tetralone, acenaphthenequinone. 30 9-acetylphenanthrene, 2-acetyl-phenanthrene, 10-thioxanthenone, 3-acetyl-

29

1 phenanthrene, 3-acetylindole, 9-fluorenone, 1-indanone, 2 1,3,5-triacetylbenzene, thioxanthen-9-one, isopropylthioxanthen-9-one. 3 xanthene-9-one, 7-H-benz[de]anthracen-7-one, 1'-acetonaphthone, 4 2'-acetonaphthone, acetonaphthone, benz[de]anthracen-7-one. 5 1'-acetonaphthone, 2'-acetonaphthone, acetonaphthone, benz[a]anthracene-7,12-dione, 2,2-dimethoxy-2-phenylacetophenone, 6 7 α, α -diethoxyacetophenone, α, α -dibutoxyacetophenone, 4-benzoyl-4'-8 methyl(diphenyl sulfide) and the like. Single oxygen-generating photosensitizers such as Rose Bengal, methylene blue, and 9 10 tetraphenylporphine as well as polymeric initiators such as poly(ethylene 11 carbon monoxide) and oligo[2-hydroxy-2-methyl-1-[4-(1-12 methylvinyl)phenyl]propanone] also can be used. However, photoinitiators 13 are preferred because they generally provide faster and more efficient 14 initiation. When actinic radiation is used, photoinitiators can provide initiation 15 at longer wavelengths which are less costly to generate and present less 16 harmful side effects than shorter wavelengths. 17 18 When a photoinitiator is present, it can enhance and/or facilitate the initiation 19 of oxygen scavenging by the composition of the present invention upon 20 exposure to radiation. The amount of photoinitiator can depend on the 21 amount and type of cyclic unsaturation present in the polymer, the 22 wavelength and intensity of radiation used, the nature and amount of 23 antioxidants used, and the type of photoinitiator used. The amount of 24 photoinitiator also can depend on how the scavenging composition is used. 25 For instance, if a photoinitiator-containing composition is in a film layer, which 26 underneath another layer is somewhat opaque to the radiation used, more 27 initiator might be needed. However, the amount of photoinitiator used for

most applications ranges from about 0.01 to about 10% (By wt.) of the total

composition. Oxygen scavenging can be initiated by exposing an article

1 containing the composition of the present invention to actinic or electron 2 beam radiation, as described below. 3 4 One or more antioxidants can be incorporated into the scavenging 5 composition of the present invention to retard degradation of the components 6 during compounding and film formation. Although such additives prolong the 7 induction period for oxygen scavenging activity to occur in the absence of 8 irradiation, the layer or article (and any incorporated photoinitiator) can be 9 exposed to radiation at the time oxygen scavenging properties are required. 10 Suitable antioxidants include 2,6-di(t-butyl)-4-methylphenol(BHT), 11 2,2'-methylene-bis(6-t-butyl-p-cresol), triphenylphosphite, tris-12 (nonylphenyl)phosphite, dilaurylthiodipropionate, vitamin E (α -tocopherol), 13 octadecyl 3,5,-di-tert-butyl-4-hydroxyhydrocinnamate. 14 tetrakis[methylene(3,5-di-tert-butyl-4-hydroxyhydrocinnamate)]methane and 15 the like. 16 17 When an antioxidant is included as part of the composition of the present 18 invention, it preferably is present in an amount which prevents oxidation of the 19 components of the oxygen scavenging composition as well as other materials 20 present in a resultant blend during formation and processing; however, the 21 amount preferably is less than that which interferes with the scavenging 22 activity of the resultant layer, film, or article after initiation has occurred. The 23 amount needed in a given composition can depend on the components 24 present therein, the particular antioxidant used, the degree and amount of 25 thermal processing used to form the shaped article, and the dosage and 26 wavelength of radiation applied to initiate oxygen scavenging. Typically, such 27 antioxidant(s) are used in an amount of from about 0.01 to about 1% (by wt.). 28 29 Other additives that also can be included in the oxygen scavenging 30 composition of the present invention include, but are not necessarily limited

1 to, fillers, pigments, dyestuffs, processing aids, plasticizers, antifog agents. 2 antiblocking agents, and the like. 3 The amounts of the components used in the oxygen scavenging composition 4 5 of the present invention can affect the use and effectiveness of this 6 composition. Thus, the amounts of polymer, transition metal catalyst, and 7 any photoinitiator, antioxidant, polymeric diluents, additives, etc., can vary 8 depending on the desired article and its end use. For example, one of the 9 primary functions of the polymer described above is to react irreversibly with 10 oxygen during the scavenging process, while a primary function of the 11 transition metal catalyst is to facilitate this process. Thus, to a large extent. 12 the amount of polymer present affects the oxygen scavenging capacity of the 13 composition, i.e., the amount of oxygen that the composition can consume. 14 while the amount of transition metal catalyst affects the rate at which oxygen 15 is consumed as well as the induction period. 16 17 The composition of the present invention can provide oxygen scavenging 18 properties at a desirable rate and capacity while having good processing and 19 compatibility properties relative to compositions including conventional non-20 cyclic ethylenically unsaturated polymers. Thus, the present composition can 21 be used to provide, by itself or as a blend with diluent film-forming polymers 22 such as polyolefins and the like, a packaging material or film that can be manufactured and processed easily. Further, the subject oxygen scavenging 23 24 composition will deplete the oxygen within a package cavity without 25 substantially detracting from the color, taste, and/or odor of the product 26 contained therein. 27 28 The amount of the polymeric scavenging component contained in the subject 29 composition can range from about 1 to almost about 100%, preferably from 30 about 5 to about 97.5%, more preferably from about 10 to 95%, even more

preferably from about 15 to about 92.5%, still more preferably from about 20 1 2 to about 90%, (with all the foregoing percentages being by weight) of the 3 composition or layer made therefrom. Typically, the amount of transition 4 metal catalyst can range from 0.001 to 1% (by wt.) of the scavenging composition, based on the metal content only (i.e., excluding ligands. 5 counterions, etc.). Where one or more other scavenging compounds and/or 6 7 diluent polymers are used as part of the composition, such other materials 8 can make up as much as 99%, preferably up to about 75%, by weight of the 9 scavenging composition. Any further additives employed normally do not 10 make up more than 10%, preferably no more than about 5%, by weight of the 11 scavenging composition. 12 13 As indicated above, the composition of the present invention can be used to 14 produce a scavenging monolayer film, a scavenging layer of a multilayer film, 15 or other articles for a variety of packaging applications. Single layer articles 16 can be prepared readily by extrusion processing. Multilayer films typically are 17 prepared using coextrusion, coating, lamination or processing. Multilayer 18 films typically are prepared using coextrusion, coating, lamination or 19 extrusion/lamination as taught in, for example, U.S. Patents 5,350,622 and 20 5,529,833, the teachings of which are incorporated herein by reference as if set forth in full. At least one of the additional layers of multilayer article can 21 22 include a material having a permeance to oxygen of no more than about 5.8 x 10⁻⁸ cm³/m²•s•Pa (i.e., about 500 cm³/m²•24 hours•atm) at about 25°C. 23 24 Polymers which are commonly used in such oxygen barrier layers include 25 poly(ethylene/vinyl alcohol), poly(vinyl alcohol), polyacrylonitrile, PVC, PVDC, 26 PET, silica, and polyamides such as nylon 6, MXD6, nylon 66, as well as 27 various amide copolymers. (Metal foil layers can also provide oxygen barrier 28 properties.) Other additional layers can include on or more layers which are 29 permeable to oxygen. In one preferred packaging construction, especially 30 flexible packages for food, the layers can include (in order starting from the

1 outside of the package to the innermost layer of the package) (a) an oxygen 2 barrier layer, (b) a scavenging layer, i.e. one that includes the scavenging 3 composition described supra, and optionally, (c) an oxygen permeable layer. 4 Control of the oxygen barrier property of layer (a) provides a means to 5 regulate the scavenging life of the package by limiting the rate of oxygen 6 entry to the scavenging layer (b), thus limiting the rate of consumption of 7 scavenging capacity. Control of the oxygen permeability of layer (c) provides a means to set an upper limit on the rate of oxygen scavenging for the overall 8 9 structure independent of the composition of scavenging layer (b). This can 10 serve the purpose of extending the handling lifetime of the film in the 11 presence of air prior to sealing of the package. Furthermore, layer (c) can 12 provide a barrier to migration of the individual components or byproducts of 13 the scavenging layer into the package interior. The term "exposed to the 14 interior" refers to a portion of a packaging article having the subject 15 scavenging composition which is either directly exposed or indirectly exposed 16 (via layers which are O₂ permeable) to the interior cavity having oxygen 17 sensitive product. Even further, layer (c) also can improve the heat 18 sealability, clarity, and/or resistance to blocking of the multilayer film. Further 19 additional layers such as the layers, easy open layers, and seal layers can 20 also be used. Polymers typically used in such tie layers include, for example, 21 anhydride functional polyolefins. 22 23 The method of the present invention includes exposing the above-described 24 composition to a package cavity having an oxygen sensitive product therein. 25 A preferred embodiment provides for including a photoinitiator as part of the 26 subject composition and subjecting a film, layer, or article that includes such a 27 composition to radiation so as to initiate oxygen scavenging at desired rates. 28 The thermal radiation used in heating and processing polymers typically used 29 in packaging films (e.g., 100-250°C) advantageously does not trigger the 30 oxygen scavenging reaction.

2 The initiating radiation preferably is actinic, e.g., UV or visible light having a 3 wavelength of from about 200 to about 750 nm, preferably of from about 200 to 600 nm, and most preferably from about 200 to 400 nm. Such light can be 4 delivered in a continuous or pulsed manner. The layer, film, etc., containing 5 the oxygen scavenging composition preferably is exposed to such radiation 6 7 until it receives at least about 1 J/g of radiation, more preferably until it receives a dose in the range of about 10 to about 2000 J/g. The radiation 8 9 also can be electron-beam radiation at a dosage of at least about 2 kiloGray 10 (kG), preferably from about 10 to about 100 kG. Other potential sources of 11 radiation include ionizing radiation such as gamma, X-ray, and corona 12 discharge. Duration of exposure depends on several factors including, but 13 not limited to, the amount and type of photoinitiator present, thickness of the 14 layers to be exposed, thickness and opacity of intervening layers, amount of 15 any antioxidant present, and the wavelength and intensity of the radiation 16 source. 17 When using oxygen scavenging layers or articles, irradiation can occur during 18 19 or after the layer or article is prepared. If the resulting layer or articles is to be 20 used to package an oxygen sensitive product, exposure can be just prior to, 21 during, or after packaging. For best uniformity of radiation, exposure 22 preferably occurs at a processing stage where the layer or article is in the 23 form of a flat sheet. For further information on initiation via irradiation, the 24 reader is directed to PCT publications WO 98/05555 and WO 98/05703, as 25 well as PCT 97/13598, 97/13370, 97/13369, the teachings of which are 26 incorporated herein by reference. 27 28 Determining the oxygen scavenging rate and capacity of a given oxygen 29 scavenging composition contemplated for a particular use can be beneficial. 30 To determine the rate, the time elapsed before the scavenger depletes a

certain amount of oxygen from a sealed container is measured. In some 1 instances, the rate can be determined adequately by placing a film containing 2 3 the desired scavenging composition in an air-tight, sealed container of an oxygen containing atmosphere, e.g., air which typically contains 20.6% (by 4 5 vol.) O₂. Over time, samples of the atmosphere inside the container are removed to determine the percentage of oxygen remaining. (Usually, the 6 7 specific rates obtained vary under different temperature and atmospheric 8 conditions. Atmospheres having lower initial oxygen content and/or 9 maintained under low temperature conditions provide a more stringent test of the scavenging ability and rate of a composition. The rates which follow are 10 11 at room temperature and one atmosphere of air, unless otherwise specified.) 12 When an active oxygen barrier is needed, a useful scavenging rate can be as low as about 0.05 cm³ oxygen per gram of the polymer in the scavenging 13 14 composition per day in air at 25°C and at 1 atm (101.3 kPa). However, in most instances, the present composition has a rate equal to or greater than 15 about 5.8 x 10⁻⁸ cm³/g•s(0.5 cm³/g•day), even up to or greater than about 16 17 5.8 x 10⁻⁵ cm³/g•s (5 cm³/g•day). Further, films or layers including the subject 18 composition are capable of a scavenging rate greater than about 1.2 x 10⁻⁴ cm³/m²•s (10 cm³/m²•day) and under some conditions, greater than 19 20 about 2.9 x 10⁻⁴ cm³/m²•s (25 cm³/m²•day). (Generally, films or layers 21 generally deemed suitable for use as an active oxygen barrier can have a 22 scavenging rate as low as 1.2 x 10⁵ cm³/m²•s (1 cm³/m²•day) when measured 23 in air at 25°C and 101 kPa (1 atm). Such rates make those layers suitable for 24 scavenging oxygen from within a package, as well as suitable for active oxygen barrier applications. 25 26 27 When the method of the present invention is to be used in an active oxygen barrier application, the initiated oxygen scavenging activity, in combination 28 29 with any oxygen barriers, preferably creates an overall oxygen permeance of less than about 1.1 x 10⁻¹⁰ cm³/m²•s•Pa (1.0 cm³/m²•day•atm) at 25°C. The 30

1 oxygen scavenging capacity preferably is such that this value is not exceeded 2. for at least two days. 3 4 Once scavenging has been initiated, the scavenging composition, layer, or 5 article prepared therefrom preferably is able to scavenge up to its capacity. 6 i.e., the amount of oxygen which the scavenger is capable of consuming 7 before it becomes ineffective. In actual use, the capacity required for a given 8 application can depend on the quantity of oxygen initially present in the 9 package, the rate of oxygen entry into the package in the absence of the 10 scavenging property, and the intended shelf life for the package. When using 11 scavengers that include the composition of the present invention, the capacity 12 can be as low as 1 cm³/g, but can be 50 cm³/g or higher. When such 13 scavengers are in a layer of a film, the layer preferably has an oxygen capacity of at least about 9.8 cm³/m² per um thickness (1200 cm³/m² per mil). 14 15 16 The composition of the present invention has been found to be capable of 17 providing a film, layer or article which substantially retains its physical 18 properties (e.g., tensile strength and modulus) even after substantial oxygen 19 scavenging has occurred. In addition, the present composition does not 20 provide significant amounts of byproducts and/or effluents, which can impart 21 an undesired taste, color, and/or odor to the packaged product. 22 23 This invention relates to an oxygen scavenging polymer composition 24 comprising cyclic allylic pendent groups which can be used in oxygen 25 scavenging packaging material which have either no or low volatile oxidation 26 by-products. Minimizing volatile by-products reduces the problem of 27 organoleptics in oxygen scavenging food packaging. 28 The polymer composition with cyclic allylic pendent groups can be made by 29 grafting methyl cyclohex-1-ene-4-methanol, cyclohex-1-ene-4-methanol

- 1 (1,2,5,6-tetrahydrobenzyl alcohol) and cyclohex-1-ene-4-propanol onto EMAC
- 2 resins by transesterification of the corresponding alcohols or transamidation
- 3 of the corresponding amines with the methyl esters on EMAC to give modified
- 4 EMAC having pendent cyclic olefins (see Figure 1). The composition can
- 5 also be made by direct polymerization.
- 6 The esterification, transesterification, amidation or transamidation reaction
- 7 can be a solution reaction or by reactive extrusion. The catalysts can be any
- 8 one of strong non-oxidizing acids, tertiary amines, Group I alkoxides,
- 9 Group IVB alkoxides and Group IVA metal organics. The level of olefin in the
- 10 final products can be controlled by the level of transesterification and the
- 11 methyl ester content of the start EMAC. The molecular weight of the
- 12 polymers largely depends on the molecular weight of the EMAC feeds.
- 13 In a preferred embodiment, these products are combined with a transition-
- 14 metal salt to catalyze the oxygen scavenging properties of the materials. A
- 15 transition-metal salt, as the term is used here, comprises an element chosen
- 16 from the first, second and third transition series of the periodic table of the
- 17 elements, particularly one that is capable of promoting oxidation reactions.
- 18 This transition-metal salt is in a form which facilitates or imparts scavenging of
- 19 oxygen by the composition of this invention. A plausible mechanism, not
- 20 intended to place limitations on this invention, is that the transition element
- 21 can readily inter-convert between at least two oxidation states and facilitates
- 22 formation of free radicals. Suitable transition-metal elements include, but are
- 23 not limited to, manganese II or III, iron II or III, cobalt II or III, nickel II or III,
- copper I or II, rhodium II, III or IV, and ruthenium. The oxidation state of the
- 25 transition-metal element when introduced into the composition is not
- 26 necessarily that of the active form. It is only necessary to have the transition-
- 27 metal element in its active form at or shortly before the time that the
- 28 composition is required to scavenge oxygen. The transition-metal element is

- 1 preferably iron, nickel or copper, more preferably manganese and most
- 2 preferably cobalt.
- 3 Suitable counter-ions for the transition metal element are organic or inorganic
- 4 anions. These include, but are not limited to, chloride, acetate, stearate,
- 5 oleate, palmitate, 2-ethylhexanoate, citrate, glycolate, benzoate,
- 6 neodecanoate or naphthenate. Organic anions are preferred. Particularly
- 7 preferable salts include cobalt 2-ethylhexanoate, cobalt benzoate, cobalt
- 8 stearate, cobalt oleate and cobalt neodecanoate. The transition-metal
- 9 element may also be introduced as an ionomer, in which case a polymeric
- 10 counter-ion is employed.
- 11 The composition of the present invention when used in forming a oxygen
- 12 scavenging packaging article can be composed solely of the above described
- polymer and transition metal catalyst. However, components, such as
- 14 photoinitiators, can be added to further facilitate and control the initiation of
- 15 oxygen scavenging properties. For instance, it is often preferable to add a
- 16 photoinitiator, or a blend of different photoinitiators, to the oxygen scavenger
- 17 compositions, especially when antioxidants are included to prevent premature
- 18 oxidation of that composition during processing.
- 19 Suitable photoinitiators are well known in the art. Such photoinitiators are
- 20 discussed in U.S. Patent No. 5,211,875. It is also discussed in U.S. Patent
- 21 Application Serial No. 08/857,325, in which some of the present inventors
- were contributing inventors and which is incorporated herein by reference.
- 23 Specific examples include, but are not limited to, benzophenone, o-methoxy-
- 24 benzophenone, acetophenone, o-methoxy-acetophenone,
- acenaphthenequinone, methyl ethyl ketone, valerophenone, hexanophenone,
- 26 α-phenyl-butyrophenone, p-morpholinopropiophenone, dibenzosuberone,
- 27 4-morpholinobenzophenone, benzoin, benzoin methyl ether,
- 28 4-o-morpholinodeoxybenzoin, p-diacetylbenzene, 4-aminobenzophenone,

- 1 4'-methoxyacetophenone, substituted and unsubstituted anthraquinones,
- 2 α-tetralone, 9-acetylphenanthrene, 2-acetyl-phenanthrene,
- 3 10-thioxanthenone, 3-acetyl-phenanthrene, 3-acetylindole, 9-fluorenone,
- 4 1-indanone, 1,3,5-triacetylbenzene, thioxanthen-9-one, xanthene-9-one,
- 5 7-H-benz[de]anthracen-7-one, benzoin tetrahydropyranyl ether.
- 6 4,4'-bis(dimethylamino)-benzophenone, 1'-acetonaphthone,
- 7 2'-acetonaphthone, acetonaphthone and 2,3-butanedione,
- 8 benz[a]anthracene-7,12-dione, 2,2-dimethoxy-2-phenylacetophenone,
- 9 α, α -diethoxy-acetophenone, α, α -dibutoxyacetophenone, etc. Singlet oxygen
- 10 generating photosensitizers such as Rose Bengal, methylene blue, and
- 11 tetraphenyl porphine may also be employed as photoinitiators. Polymeric
- 12 initiators include polyethylene carbon monoxide and oligo[2-hydroxy-2-
- methyl-1-[4-(1-methylvinyl)phenyl]propanone]. Use of a photoinitiator is
- 14 preferable because it generally provides faster and more efficient initiation.
- 15 When a photoinitiator is used, its primary function is to enhance and facilitate
- the initiation of oxygen scavenging upon exposure to radiation. The amount
- 17 of photoinitiator can vary. In many instances, the amount will depend on the
- 18 amount and type of oxygen scavenging polymer in the present invention, the
- 19 wavelength and intensity of radiation used, the nature and amount of
- antioxidants used, as well as the type of photoinitiator used. The amount of
- 21 photoinitiator also depends on how the scavenging composition is used. For
- 22 instance, if the photoinitiator-coating composition is placed underneath a layer
- 23 which is somewhat opaque to the radiation used, more initiator may be
- 24 needed. For most purposes, however, the amount of photoinitiator, when
- used, will be in the range of 0.01 to 10% by weight of the total composition.
- 26 The initiating of oxygen scavenging can be accomplished by exposing the
- 27 packaging article to actinic or electron beam radiation, as described below.

- 1 Antioxidants may be incorporated into the scavenging compositions of this
- 2 invention to control degradation of the components during compounding and
- 3 shaping. An antioxidant, as defined herein, is any material which inhibits
- 4 oxidative degradation or cross-linking of polymers. Typically, such
- 5 antioxidants are added to facilitate the processing of polymeric materials
- 6 and/or prolong their useful shelf-life.
- 7 Antioxidants such as Vitamin E, Irganox® 1010, Irganox® 1076,
- 8 2,6-di(t-butyl)-4-methyl-phenol(BHT), 2,6-di(t-butyl)-4-ethyl-phenol (BHEB),
- 9 2,2'-methylene-bis(6-t-butyl-p-cresol), triphenylphosphite,
- 10 tris-(nonylphenyl)phosphite and dilaurylthiodipropionate would be suitable for
- 11 use with this invention.
- When an antioxidant is included as part of the present composition, it should
- be used in amounts which will prevent oxidation of the scavenger
- 14 composition's components as well as other materials present in a resultant
- 15 blend during formation and processing but the amount should be less than
- that which would interfere with the scavenging activity of the resultant layer,
- 17 film or article. The particular amount needed will depend on the particular
- 18 components of the composition, the particular antioxidant used, the degree
- 19 and amount of thermal processing used to form the shaped article, and the
- 20 dosage and wavelength of radiation applied to initiate oxygen scavenging and
- 21 can be determined by conventional means. Typically, they are present in
- 22 about 0.01 to 1% by weight.
- 23 Other additives which may also be included in oxygen scavenger layers
- include, but are not necessarily limited to, fillers, pigments, dyestuffs,
- 25 stabilizers, processing aids, plasticizers, fire retardants, anti-fog agents, etc.
- 26 The amounts of the components which are used in the oxygen scavenging
- 27 compositions, or layers have an effect on the use, effectiveness and results of

- 1 this method. Thus, the amounts of polymer, transition metal catalyst and any
- 2 photoinitiator, antioxidant, polymeric diluents and additives, can vary
- 3 depending on the article and its end use.
- 4 For instance, one of the primary functions of the polymer described above is
- 5 to react irreversibly with oxygen during the scavenging process, while the
- 6 primary function of the transition metal catalyst is to facilitate this process.
- 7 Thus, to a large extent, the amount of polymer present will affect the oxygen
- 8 scavenging capacity of the composition, i.e., affect the amount of oxygen that
- 9 the composition can consume. The amount of transition metal catalyst will
- affect the rate at which oxygen is consumed. Because it primarily affects the
- 11 scavenging rate, the amount of transition metal catalyst may also affect the
- 12 onset of oxygen scavenging (induction period).
- 13 It has been found that the subject polymers, when used as part of the present
- 14 composition, provide oxygen scavenger properties at desirable rate and
- 15 capacity while causing the composition to have enhanced processability and
- 16 compatibility properties over conventional ethylenically unsaturated polymers.
- 17 Thus, the present composition can be used to provide, by itself or as a blend
- with diluent polymers, such as polyolefins and the like, a packaging material
- 19 or film having enhanced processability properties. Further, the present
- 20 composition consumes and depletes the oxygen within a package cavity
- 21 without substantially detracting from the color, taste and/or odor of the
- 22 product contained within the package cavity.
- 23 The amount of the above-described polymer contained as part of the present
- 24 composition may range from about 1 to 100% by weight of the composition or
- 25 layer composed of said composition in which both polymer and transition
- 26 metal catalyst are present (hereinafter referred to as the "scavenging
- 27 composition", e.g., in a coextruded film or container, the scavenging
- 28 composition would comprise the particular layer(s) in which both the

- 1 copolymer and transition metal catalyst components are present together).
- 2 Typically, the amount of transition metal catalyst may range from 0.001 to 1%
- 3 (10 to 10,000 ppm) of the scavenging composition, based on the metal
- 4 content only (excluding ligands, counterions, etc.). In the event the amount of
- 5 transition metal catalyst is less than 1%, it follows that the polymer and any
- 6 additives will comprise substantially all of the remainder of the composition.
- 7 The polymer of the present invention may further be combined with other
- 8 polymeric oxygen scavenger agents.
- 9 Any further additives employed normally will not comprise more than 10% of
- 10 the scavenging composition, with preferable amounts being less than 5% by
- weight of the scavenging composition.
- 12 Optionally, the compositions and process of this invention can include
- 13 exposure of the polymer containing the oxygen scavenging-promoting
- transition metal to actinic radiation to reduce the induction period, if any,
- 15 before oxygen scavenging commences. A method is known for initiating
- 16 oxygen scavenging by exposing a film comprising an oxidizable organic
- 17 compound and a transition metal catalyst to actinic radiation. A composition
- of the present invention which has a long induction period in the absence of
- 19 actinic radiation but a short or non-existent induction period after exposure to
- 20 actinic radiation is particularly preferred. They maintain a high capability for
- 21 scavenging oxygen upon activation with actinic radiation. Thus, oxygen
- 22 scavenging can be activated when desired.
- 23 The radiation used in this method should be actinic, e.g., ultraviolet or visible
- 24 light having a wavelength of about 200 to 750 nanometers (nm), and
- 25 preferably having a wavelength of about 200 to 600 nm, and most preferably
- 26 from about 200 to 400 nm. When employing this method, it is preferable to
- 27 expose the oxygen scavenger to at least 0.01 Joule per gram of scavenging
- 28 composition. A typical amount of exposure is in the range of 10 to

1 2000 Joules per gram. The radiation can also be an electron beam radiation 2 at a dosage of about 2 to 200 kiloGray, preferably about 10 to 100 kiloGray. 3 Other sources of radiation include ionizing radiation such as gamma, X-rays 4 and corona discharge. The duration of exposure depends on several factors including, but not limited to, the amount and type of photoinitiator present, 5 6 thickness of the layers to be exposed, thickness and opacity of intervening 7 layers amount of any antioxidant present, and the wavelength and intensity of 8 the radiation source. The radiation provided by heating of polyolefin and the 9 like polymers (e.g., 100-250°C) during processing does not cause triggering. 10 Oxygen-scavenging compositions of the present invention are useful in many 11 ways. The compositions can be dispersed as small particles for absorbing 12 oxygen or can be coated onto materials such as metallic foil, polymer film. 13 metalized film, paper or cardboard to provide, in some embodiments, 14 scavenging properties and/or adhesive properties. The compositions are also 15 useful in making articles such as single or multi-layer rigid thick-walled plastic 16 containers or bottles (typically, between 5 and 100 mils in thickness) or in 17 making single or multi-layer flexible films, especially thin films (less than 18 5 mils, or even as thin as about 0.25 mil). Some of the compositions of the 19 present invention are easily formed into films using well-known means. 20 These films can be used alone or in combination with other films or materials. 21 The compositions of the present invention may be further combined with one 22 or more polymers, such as thermoplastic polymers which are typically used to 23 form film layers in plastic packaging articles. In the manufacture of certain 24 packaging articles, well-known thermosets can also be used as a polymeric 25 diluent. 26 Selecting combinations of a diluent and the composition of the present 27 invention depends on the properties desired. Polymers which can be used as

the diluent include, but are not limited to, polyethylene, low or very low density

- polyethylene, polypropylene, polyvinyl chloride, and ethylene copolymers such as ethylene-vinyl acetate, ethylene-alkyl acrylates or methacrylates,
- 3 ethylene-acrylic acid or methacrylic acid, and ethylene-arylic or methacrylic
- 4 acid ionomers. In rigid packaging applications, polystyrene is often used.
- 5 Blends of different diluents may also be used. However, as indicated above,
- 6 the selection of the polymeric diluent largely depends on the article to be
- 7 manufactured and the end use. Such selection factors are well known in the
- 8 art.
- 9 If a diluent polymer such as a thermoplastic is employed, it should further be
- selected according to its compatibility with the composition of the present
- 11 invention. In some instances, the clarity, cleanliness, effectiveness as an
- 12 oxygen-scavenger, barrier properties, mechanical properties and/or texture of
- 13 the article can be adversely affected by a blend containing a polymer which is
- 14 incompatible with the composition of the present invention.
- 15 A blend of a composition of the present invention with a compatible polymer
- 16 can be made by dry blending or by melt-blending the polymers together at a
- 17 temperature in the approximate range of 50°C to 250°C. Alternative methods
- of blending include the use of a solvent followed by evaporation. When
- 19 making film layers or articles from oxygen-scavenging compositions, extrusion
- 20 or coextrusion, solvent casting, injection molding, stretch blow molding.
- orientation, thermoforming, extrusion coating, coating and curing, lamination
- 22 or combinations thereof would typically follow the blending.
- 23 Layers comprising the composition of the present invention may be in several
- forms. They may be in the form of stock films, including "oriented" or "heat
- shrinkable" films, which may ultimately be processed as bags, etc., or in the
- 26 form of stretch-wrap films. The layers may also be in the form of sheet inserts
- 27 to be placed in a packaging cavity. In rigid articles such as beverage
- 28 containers, thermoformed trays or cups, the layer may be within the

- 1 container's walls. Even further, the layer may also be in the form of a liner
- 2 placed with or in the container's lid or cap. The layer may even be coated or
- 3 laminated onto any one of the articles mentioned above.
- 4 In multi-layered articles, the scavenging layer comprising the composition of
- 5 the present invention may be included with layers such as, but not necessarily
- 6 limited to, "oxygen barriers", i.e., layers of material having an oxygen
- 7 transmission rate equal to or less than 100 cubic centimeters-mil per square
- 8 meter (cc-mil/m²) per day per atmosphere pressure at room temperature, i.e.,
- 9 about 25°C. Typical oxygen barriers comprise poly(ethylene vinyl alcohol),
- 10 polyacrylonitrile, polyvinyl chloride, poly(vinylidene dichloride), polyethylene
- 11 terephthalate, silica and polyamides. Metal foil layers can also be employed.
- 12 Other additional layers may include one or more layers which are permeable
- 13 to oxygen. In one preferred packaging construction, especially for flexible
- 14 packaging for food, the layers include, in order starting from the outside of the
- package to the innermost layer of the package, (i) an oxygen barrier layer,
- 16 (ii) a scavenging layer, i.e., the scavenging composition as defined earlier,
- 17 and, optionally, (iii) an oxygen permeable layer. Control of the oxygen barrier
- 18 property of (i) allows a means to regulate the scavenging life of the package
- 19 by limiting the rate of oxygen entry to the scavenging composition (ii), and
- 20 thus limiting the rate of consumption of scavenging capacity. Control of the
- 21 oxygen permeability of layer (iii) allows a means to set an upper limit on the
- 22 rate of oxygen scavenging for the overall structure independent of the
- 23 composition of the scavenging composition (ii). This can serve the purpose of
- 24 extending the handling lifetime of the films in the presence of air prior to
- 25 sealing of the package. Furthermore, layer (iii) can provide a barrier to

ı

- 26 migration of the individual components in the scavenging films or by-products
- 27 of scavenging into the package interior. Even further, layer (iii) also improves
- 28 the heat-sealability, clarity and/or resistance to blocking of the multi-layer film.

- 1 Further, additional layers such as adhesive layers may also be used.
- 2 Compositions typically used for adhesive layers include anhydride functional
- 3 polyolefins and other well-known adhesive layers.
- 4 To determine the oxygen scavenging capabilities of a composition, the rate of
- 5 oxygen scavenging can be calculated by measuring the time that elapsed
- 6 before the article depletes a certain amount of oxygen from a sealed
- 7 container. For instance, a film comprising the scavenging component can be
- 8 placed in an air-tight, sealed container of a certain oxygen containing
- 9 atmosphere, e.g., air which typically contains 20.9% oxygen by volume.
- 10 Then, over a period of time, samples of the atmosphere inside the container
- 11 are removed to determine the percentage of oxygen remaining. The
- scavenging rates of the composition and layers of the present invention will
- 13 change with changing temperature and atmospheric conditions.
- 14 When an active oxygen barrier is prepared, the scavenging rate can be as
- 15 low as 0.1 cc oxygen per gram of composition of the present invention per
- day in air at 25°C and a 1 atmosphere pressure. However, preferable
- 17 compositions of this invention have rates equal to or greater than 1 cc oxygen
- per gram per day, thus making them suitable for scavenging oxygen from
- 19 within a package, as well as suitable for active oxygen barrier applications.
- 20 Many compositions are even capable of more preferable rates equal to or
- 21 greater than 5.0 cc O₂ per gram per day.
- 22 Generally, film layers suitable for use as an active oxygen barrier can have an
- 23 oxygen transmission rate as high as 10 cc oxygen per square meter per mil
- 24 per day when measured in air at 25°C and 1 atmosphere pressure.
- 25 Preferably, a layer of this invention has an oxygen transmission rate less than
- about 1 cc oxygen per square meter per mil per day, and more preferably has
- 27 an oxygen transmission rate less than about 0.2 cc oxygen per square meter
- per rail per day under the same conditions, thus making it suitable for active

- 1 oxygen barrier applications as well as for scavenging oxygen from within a
- 2 package.
- 3 In an active oxygen barrier application, it is preferable that the combination of
- 4 oxygen barriers and any oxygen scavenging activity create an overall oxygen
- 5 transmission rate of less than about 1.0 cubic centimeter-mil per square
- 6 meter per day per atmosphere pressure at 25°C. Another definition of
- 7 acceptable oxygen scavenging is derived from testing actual packages. In
- 8 actual use, the scavenging rate requirement will largely depend on the
- 9 internal atmosphere of the package, the contents of the package and the
- 10 temperature at which it is stored.
- 11 In a packaging article made according to this invention, the scavenging rate
- will depend primarily on the amount and nature of the composition of the
- present invention in the article, and secondarily on the amount and nature of
- other additives (e.g., diluent polymer, antioxidant, etc.) which are present in
- 15 the scavenging component, as well as the overall manner in which the
- package is fabricated, e.g., surface area/volume ratio.
- 17 The oxygen scavenging capacity of an article comprising the invention can be
- 18 measured by determining the amount of oxygen consumed until the article
- 19 becomes ineffective as a scavenger. The scavenging capacity of the
- 20 package will depend primarily on the amount and nature of the scavenging
- 21 moieties present in the article, as discussed above.
- 22 In actual use, the oxygen scavenging capacity requirement of the article will
- 23 largely depend on three parameters of each application:
- 24 (1) the quantity of oxygen initially present in the package;

1	(2) the rate of oxygen entry into the package in the absence of the
2	scavenging property; and
3	(3) the intended shelf life for the package.
4	The scavenging capacity of the composition can be as low as 1 cc oxygen per
5	gram, but is preferably at least 10 cc oxygen per gram, and more preferably
6	at least 50 cc oxygen per gram. When such compositions are in a layer, the
7	layer will preferably have an oxygen capacity of at least 250 cc oxygen per
8	square meter per mil thickness and more preferably at least 500 cc oxygen
9	per square meter per mil thickness.
10	Other factors may also affect oxygen scavenging and should be considered
11	when selecting compositions. These factors include but are not limited to
12	temperature, relative humidity, and the atmospheric environment in the
13	package.
14	Applicants have achieved a composition for a rigid beverage and food
15	container comprising PET and/or PEN, the container incorporating an oxygen
16	scavenging component of cyclic olefin which oxidizes oxygen in the interior of
17	the container without giving off odor and/or taste as a result of its oxygen
18	scavenging function, nor does it cause a change in molecular weight. This is
19	because the cyclic olefin oxygen scavenging component does not fragment
20	as it oxidizes, Thus the composition maintains the structural integrity of the
21	container while avoiding the problem of imparting oxidation byproducts to the
22	packaged material.
23	
24	EXAMPLES
25	Objects and advantages of this invention are further illustrated by the
26	following examples. The particular materials and amounts thereof, as well as

Example 4

1	other conditions and details, recited in these examples should not be used to
2	unduly limit this invention.
3	
4	Example 1
5	Preferred embodiments of the present invention include polymers and
6	oligomers, which contain cyclohexene groups accessible to free oxygen
7	molecules. These polymers or oligomers may be prepared from any of a
8	number of methods though one preferred reaction comprises 1, 2, 3, 6,
9	tetrahydrophthalic anhydride. This anhydride is a low cost monomer derived
10	from butadiene, which makes it particularly attractive on a commercial scale.
11	The anhydride may be used to make polyester resins such as by reaction with
12	diols. It may also be reacted with hydroxy or polyhydroxy compounds to
13	produce half esters suitable for subsequent use in plastic film and materials
14	manufacture.
15	
16	Example 2
17	Non-aromatic alkenyl benzyl alcohols (e.g. tetrahydrobenzyl alcohols) may
18	also be reacted with certain compounds to produce useful scavengers. For
19	instance tetrahydrobenzyl alcohol may be reacted with compounds containing
20	a carboxylic acid, acid halide, ester, anhydride and/or isocyanate functionality
21	These compounds may be small molecules or oligomers or polymers. For
22	example, tetrahydrobenzyl alcohol may be reacted with styrene, maleic
23	anhydride copolymers or with polyfunctional isocyanates.
24	
25	Example 3
26	Cyclohexene dimethanol compounds may be used to prepare oxygen
27	absorbing polyesters and polyurethanes.
28	•

1	As another example, tetrahydrobenzoic acid and tetrahydrobenzaldehyde
2	may also be used to modify various hydroxyl functional materials.
3	
4	Example 5
5	Reactions such as the functionalization of polymers may be carried out by a
6	reactive extrusion process. For instance this may be a transesterification
7	process.
8	
9	Example 6
10	Cyclohexene anhydride may be used in the preparation of useful oxygen
11	scavengers. These cyclohexene anhydrides may be prepared by from a
12	diene monomer such as butadiene with maleic anhydride. Of commercial
13	attractiveness are their low cost and their ability to be converted into a
14	number of useful intermediates. In addition, they may also be used to
15	functionalize OH containing polymers. The half esters, which form rapidly
16	when a cyclic anhydride reacts with an OH group, may be subsequently
17	neutralized and the resultant materials dispersed in ionomers or ethylene
18	acrylic acid copolymers (for instance).
19	· ·
20	For ease of use, small functionalized molecules such as the reaction product
21	of four moles of tetrahydrophthalic anhydride with pentaerythritol may be
22	prepared either by heating in a mutual solvent or by a reactive extrusion
23	process. These may then be dispersed into a commodity polymer such as
24	EVA.
25	•
26	The cyclohexene anhydrides may also be converted into linear polyesters by
27	reaction with ethylene glycol and the like.
28	
29	Example 7

1 Useful anhydrides are cyclic anhydrides and in particular the Diels Alder 2 adducts of various alkenes. Typically this will comprise 1, 3 butadiene (and 3 substituted derivatives) with other compounds able to complete a Diels Alder 4 type reaction. The resulting anhydrides may then be used in the manufacture 5 of various oxygen scavengers, and polymers containing same. 6 7 ′ Example 8 8 Film structures, coatings, and molded articles, as well as sachets and 9 impregnated matrices, are envisaged incorporating oxygen scavengers as 10 discussed previously. Also included are transition metal catalysts such as 11 used in the prior art for catalyzing oxygen scavenging reactions. Optionally 12 initiators or triggers for the reaction may also be included. 13 14 Example 9 15 16 Preparation of a low molecular weight oxidizable oil from 3-Cyclohexene-1-17 carbonyl chloride and triethylene glycol. 18 19 3-Cyclohexene-1-carbonyl chloride was prepared as follows: 20 21 50g of thionyl chloride was added to 27.6g of 3-cyclohexene-1-carboxylic acid 22 and the solution was stirred for two hours at 50°C. Excess thionyl chloride 23 was removed under vacuum and the resulting yellow brown oil was purified 24 by distillation under vacuum (bp 80-82°C at 18-19mm Hg). 25 26 The oil was then prepared in the following manner: 27 28 In a 250 ml flask fitted with a drying tube was placed 18.7g of 3-cyclohexene-29 1-carbonyl chloride and 40cc of methylene chloride. A solution of 9.6g of 30 triethylene glycol in 20ml of methylene chloride was added and the reaction

1	was stirred for 2 hours at room temperature, by which time the evolution of		
2	hydrochloric acid had ceased.		
3			
4	80ml of 10% aqueous sodium bicarbonate was added to the reaction mixture		
5	and the mixture was vigorously stirred for 45 minutes. The organic layer was		layer was
6	collected, washed with water and then dried with magnes	sium sulph	ate. The
7	methylene chloride was removed under reduced pressure giving a colorless		colorless
8	oil.		
9	The cyclohexene oil was compounded into a film with the	following	parts by
10	weight:	•	
11			
12	Oil	12	2
13	Silica	5	
14	Benzophenone	0.3	3
15	Cobalt (111) acetylacetonate	0.28	
16	Ethylene vinyl acetate copolymer (18% EVA)	90	
17			
18	A similar film was prepared using sunflower seed oil in place of the		
19	cyclohexene based oil.		
20			
21	Both films were exposed to 4 minutes of UV light, then sealed in oxygen		ygen
22	barrier bags and stored in the dark.		
23			
24	Both materials scavenged oxygen after photoexposure and the sunflower oil		flower oil
25	based material was a faster scavenger than the cyclohex	ene oil ba	sed
26	material. However, gas chromatography of the headspace	ce of the b	ags post
27	oxidation revealed that there was a very large difference	in the leve	ls of
28	volatile components. The cyclohexene based material pr	roduced le	ss than
29	3% of the volatile components produced by the sunflower oil based material.		material.

1	
2	The cyclohexene based films were stable for more than 300 days if stored at
3	room temperature in the absence of light (i.e., the oxygen concentration in a
4	sealed package containing the film specimens was essentially unchanged
5	after storage for this time period).
6	
7	A similar cyclohexene based film was prepared, this time using 3,4 dimethyl-
8	3-cyclohexene-1-carbonyl chloride as the starting material. This film was a
9	much faster oxygen absorber than the film prepared from the unsubstituted
10	product. The film form the substituted produced less than 10% of the total
11	volatile components produced from an equivalent film made from sunflower
12	oil.
13	
14	The dimethyl cyclohexene based films were stable for at least two hundred
15	days when stored at room temperature in the absence of light. The stability of
16	similar vegetable oil based films was limited to around 50 days.
17	
18	This series of experiments revealed the following:
19	
20	1. Cyclohexene functionalized materials are effective oxygen absorbers.
21	2. The speed of reaction may be increased by substituting methyl groups
22	adjacent to the double bond.
23	3. Cyclic alkene based materials produce much lower levels of volatile
24	oxidation products than linear alkene based materials.
25	4. The storage stability of cyclohexene containing films is excellent.
26	
27	Example 10
28	Preparation of an oxidizable polyester resin.

1	In a three neck round bottom flask equipped with a Dean and Stark trap,	
2	reflux condenser and nitrogen inlet/exit were placed the following materials:	
3		
4	cis-1,2,3,6-Tetrahydrophthalic anhydride 35.54g	
5	1,4-Butanediol 20g	
6		
7	75ml of xylene was added, so that the trap was full of xylene and the mixture	
8	was brought to reflux. The reaction was refluxed for six and a half hours:	
9	0.55g of p-Toluenesulfonic acid monohydrate was added and reflux was	
10	continued for a further six and a half hours.	
11	25ml of xylene was removed from the trap and the mixture was refluxed for a	
12	further one hour. A very viscous pale colored solution was obtained.	
13		
14	The solution was extracted with methanol to remove the acid catalyst, and	
15	was diluted with dichloromethane prior to use.	
16		
17	The polymer was obtained as a 38% w/w solids solution in	
18	toluene/dichloromethane. To 12.37 g of the polymer solution was taken	
19	0.0213 g cobalt Ten-Cem® (OMG Inc.) in 5 mL of dichloromethane and	
20	0.0069 g of Quantacure™CPTX (1-chloro-4-propoxy-thioxanthone, Great	
21	Lakes Fine Chemicals) was added. The mixture was stirred for a few minutes	
22	and a film was cast onto the surface of the another film at a wet thickness of	
23	about 1 mm. A second film was formulated as follows and cast as above:	
24	12.64 g polymer solution, 0.0318 g cobalt Ten-Cem® and 0.0074 g	
25	4,4'-dimethoxybenzophenone (DMOBP, Spectrum Quality Products Inc.).	
26		
27	The dried films were irradiated for 2.5 minutes with a combination of	
28	germicidal and backlight UV lamps. The approximate dose of UVC was	
29	1350mJcm ² and the approximate dose of UVA was 1950mJcm ² . The	

- 1 irradiated films were sealed in a barrier pouch along with 120cc of air. The
- 2 oxygen content was monitored with time and the following results were

3 obtained:

Film #1, 4.7g, 1020 ppm Cobalt and 1470 ppm CPTX	
Percent Oxygen	
20.6	
19.2	
18.4	
16.9	

4

Film #2, 4.8g, 1500 ppm Cobalt and 1480 ppm DMOBP	
Elapsed Time (days)	Percent Oxygen
0	20.6
1	19.3
2	18.6
3	17.0

5

- 6 This example illustrates that polyesters derived from tetrahydrophthalic
- 7 anhydride are useful oxygen scavengers.

8

- Example 11
- 10 Preparation of an oxidizable polymer from 3-Cyclohexene-1-methanol and an
- 11 alternating copolymer of maleic anhydride and octadecene.

- 13 In a three neck round bottom flask equipped with condenser and nitrogen
- inlet was placed 20 g of poly(maleic anhydride-alt-1-octadecene). 80 cc of
- 15 methylene chloride was added and the mixture was stirred to dissolve. After

1 a clear solution had been obtained 3.2g of 3-Cyclohexene-1-methanol was 2 added, and washed into the flask with a further 10cc of methylene chloride. 3 4 The mixture was refluxed with stirring under nitrogen for two hours, then left 5 overnight at room temperature. The solution was refluxed for a further three 6 hours and allowed to cool to room temperature. 7 8 The polymer was obtained as a 21.9 wt. % solution in dichloromethane. To 9 20.51 g of the polymer solution was added 0.0201 g of cobalt Ten-Cem® 10 (OMG Inc., 22.5% Co by wt.) dissolved in 5 mL of toluene solution and 0.0038 g of Quantacure™ BMS (4-benzoyl-4'-methyl(diphenyl sulfide) 11 12 available from Great Lakes Fine Chemicals Ltd.). The mixture was stirred for 13 a few minutes and a film was cast using a draw down bar to a wet film 14 thickness of about 1 mm. 15 16 A second film was formulated as follows: 20.10 g polymer solution, 0.0474 g 17 cobalt Ten-Cem®, 0.0079 g 4,4'-dimethylbenzophenone (DMBP, from 18 Lancaster Synthesis). A third film was formulated as follows: 20.84 g 19 polymer solution, 0.0398 g cobalt Ten-Cem®, 0.0085 g 20 2-isopropylthioxanthone (ITX, First Chemical Co.). 21 22 The dried films were irradiated for 2.5 minutes with a combination of 23 germicidal and backlight UV lamps. The approximate dose of UVC was 1350 mJ/cm² and the approximate dose of UVA was 1950 mJ/cm². The 24 25 irradiated films were sealed in a barrier pouch along with about 120 cc of air. 26 The oxygen content was monitored with time as described elsewhere. The 27 following results were obtained: 28

Film Sample #1, 1.34g, with 1004 ppm Cobalt and 844 ppm BMS		
Elapsed Time (days)	Percent Oxygen	
0	20.6	
1	12.2	
2	7.5	
5	6.2	

Film Sample #2, 3.04g, with 2420 ppm Cobalt and 1795 ppm DMBP		
Elapsed Time (days)	Percent Oxygen	
0	20.6	
1	11.8	
2	10.0	
5	9.7	

2 3

Film Sample #3, 2.09g, with 1960 ppm Cobalt and 1860 ppm ITX		
Elapsed Time (days)	Percent Oxygen	
_ 0	20.6	
1	13.8	
2	10.5	
5	10.0	

4

- 5 The results suggest that the reaction of a polymeric anhydride and
- 6 tetrahydrobenzyl alcohol is a useful route to oxygen scavenging plastics.

7

- 8 Example 12
- 9 Preparation of a Cyclohexene containing polymer by transesterification

1 To a 2L resin kettle was taken 180 g of polyethylene-co-methyl acrylate 2 (EMAC® SP2260, Chevron, 24 wt. % methyl acrylate) and 1 L of toluene. The kettle was equipped with a mechanical overhead stirrer. Dean-Stark trap 3 4 and a condenser. The kettle was heated to melt the polymer. To the stirred 5 solution was added 28.12 g of 3-cyclohexene-1-methanol, followed by the addition of 2.145 g of 4-(2-hydroxyethoxy)benzophenone. (Note: this 6 7 benzophenone derivative was prepared by the method of Yoshino et al. Bull. 8 Chem Soc. Japan, 1973, 46, 553-6 using 4-hydroxybenzophenone, ethylene 9 carbonate and tetraethylammonium iodide.) The catalyst, titanium (IV) 10 isopropoxide (1.05g) was added. The mixture turned yellow and the reflux 11 rate increased. Heat was maintained for 4 hours and about 75 mL of 12 condensate was removed in four fractions. An additional 0.5 g of titanium 13 isopropoxide was added and heat was maintained for an additional 8 hours. 14 Additional toluene was added as needed to maintain the reaction volume. 15 Again an additional 0.5 g of catalyst was added and heat maintained for 16 another 8 hours. Analysis of the condensate showed no more production of 17 methanol. The reaction mixture was cooled to a gel and precipitated into 18 methanol. The polymer was washed with methanol until nothing was 19 extracted into the methanol fractions. 20 21 The above resin containing cyclohexene pendant groups and a covalently 22 bound benzophenone derivative was melt compounded with 500 ppm 23 vitamin E as the antioxidant and 10% of an EVA based cobalt (II) oleate 24 (Shepherd Chemicals) masterbatch. The masterbatch contained 1.0% cobalt metal by weight. Samples were compression molded and cut to 197.56 cm² 25 26 of UVC light (254 nm) and was sealed in an oxygen barrier pouch (Cryovac 27 P640B) with 300 cc of air and was stored in the dark at room temperature. 28 Headspace oxygen levels were monitored periodically by withdrawing a 4 cc 29 sample and analyzing using a Mocon model LC 700F oxygen analyzer. The 30 following results were obtained for the 1.9 g (7.8 mil thick) sample.

Elapsed Time (days)	Percent Oxygen
0	20.6
1	15.6
5	5.0
14	2.1
21	1.2

2

3 This example illustrates excellent oxygen scavenging ability from this type of

4 polymer and the usefulness of a covalently bound photoinitiator.

5

6 Example 13

7 Preparation of oxidizable polyurethanes.

8

9 In a two necked 250ml flask equipped with reflux condenser and nitrogen

10 inlet/exit were placed the following materials:

11

12 1,6-Diisocyanatohexane

6.5g

13 3-Cyclohexene-1,1dimethanol

5.23g

14 2-Butanone

70ml

15

- 16 One drop of dibutyltin dilaurate was added and the mixture was stirred under
- 17 nitrogen for thirty minutes at room temperature. The mixture was then
- 18 brought to reflux for a further four hours and one drop of water in 10ml of
- 19 MEK was added. The mixture was refluxed for a further hour and then
- 20 allowed to cool to room temperature.

- 22 A polyurethane containing cyclic unsaturation was prepared from 1,6-
- 23 diisocyanatohexane and 3-cyclohexene-1,1-dimethanol. The polymer
- 24 (3.912 g) was taken into 10 mL dichloromethane and a solution of 0.0243 g of

WO 99/48963

1 cobalt Ten-Cem® (OMG Inc.) in 5 mL of dichloromethane was added. To the

- 2 stirred mixture was added 0.0084 g of 4,4'-dimethylbenzophenone (DMBP,
- 3 Lancaster Synthesis). The mixture was stirred for about 15 minutes. A film
- 4 was cast from the solution on the surface of another film at a wet thickness of
 - 5 about 1 mm. The dried film was triggered and tested as described in
 - 6 example 3 above.

7

3.9g, with 1400 ppm Cobalt and 2150 ppm DMBP					
Elapsed Time (days) Percent Oxygen					
0	20.6				
1	18.3 13.1				
4					
5	9.5				

8

9 These results suggest that polyurethanes derived from 3-Cyclohexene-1,1-

10 dimethanol are useful oxygen absorbers. These materials and alternative

11 formulations may be useful in formulating oxygen scavenging adhesive resins

12 for use in flexible packaging i.e., for use in lamination.

13 14

Example 14

- 15 Preparation of a poly (vinyl acetal) from poly(vinyl alcohol) and
- 16 3-Cyclohexene-1-carboxaldehyde.

- 18 In a 500ml flask equipped with nitrogen inlet/exit and mechanical stirrer was
- 19 placed 150 ml of a 70/30 mix of dioxane/ethanol and 10g of poly(vinylalcohol).
- 20 The mixture was stirred and 15.7g of 3-Cyclohexene-1-carboxaldehyde was
- 21 added, followed by 0.25ml conc. HCl and 5 mg of hydroquinone. The mixture
- 22 was refluxed for four hours, during which time the poly(vinyl alcohol) dissolved
- 23 and turned a pale yellow color, 0.5g of sodium acetate was added followed by

- 1 2.5g of urea, both in aqueous solution. The polymer precipitated and was
- 2 purified by addition of further dioxane then precipitation into water. The dried
- 3 polymer was found to contain approximately 63mole% of acetal groups.

- 5 A similar polymer was prepared from 3,4-dimethyl-3-cyclohexene-1-
- 6 carboxaldehyde and poly(vinyl alcohol) which contained approximately
- 7 65 mole% of acetal groups.

8

- 9 A solution of cobalt (III) acetylacetonate (20mg) and benzophenenone (20mg)
- in methylene chloride was added with stirring to a solution (1g) of each acetal
- resin dissolved in 15ml of warm dioxane. The solution was poured into a
- 12 150mm diameter flat bottom petri dish and the solvent was allowed to
- 13 evaporate. The resultant film was held under high vacuum for 2-3 hours to
- 14 remove any residual solvent. A further sample containing 30% of a dibutyl
- 15 phthalate plasticiser was also prepared using the dimethyl substituted acetal
- 16 resin described above.

17

18

- The film samples were exposed to 4 minutes of UVA radiation and then
- 19 vacuum packed in a barrier bag. 200ml of air was injected into the bag and
- 20 the puncture point was isolated by heat sealing. The pouch was stored in the
- 21 absence of light.

22

The following results were obtained for the unsubstituted resin:

Elapsed Time (days)	Percent Oxygen		
0	20.6		
1	no reading		
3	9.9		
6	7.2		

12	1.2

2 The following results were obtained for the disubstituted resin:

Elapsed Time (days)	Percent Oxygen		
0	20.6		
1	10.2 4.3		
3			
. 6	1.4		
21	0		

3

4 The following results were obtained from the plasticised resin:

5

Elapsed Time (days)	Percent Oxygen		
0	20.6		
3	3.7 1.8		
4			
7	0.2		
12	0		

6

7 These results demonstrate the following principles:

8

9 1. Cyclohexene based acetal resins are effective oxygen scavengers.

10 11

2. The substituted cyclohexene rings provide faster oxygen scavengers than the unsubstituted resins.

13

12

14 3. A plasticiser tends to increase the rate of oxygen scavenging.

1	
2	
3	
4	Non-limiting examples are given in Examples 15 and 16 below of
5	experimental conditions that were used for preparation of the polymers.
6	Non-limiting examples of the resin preparation followed by steam stripping as
7	well as compounding the polymers with oxidation catalyst, such as cobalt
8	oleate and a photoinitiator, such as Methanone,
9	[5'-(5'-(4-benzoylphenyl)[1,1':3', 1"-terphenyl]-4,4"-diyl]bis[phenyl- (hereinafter
10	referred to as BBP³), and extruded into a 3-layer film having a PE/oxygen
11 .	scavenging polymer/PE structure are provided in Examples 17 through 20
12	below.
13	Headspace studies of three layer films made by compounding catalyst
14	package with both fresh and aged (20 months stored in air at ambient
15	temperature) resins after UV triggering give a very fast rate of oxygen
16	scavenging and the resulting packages are relatively non-odorous.
17	Non-limiting examples of such studies are given in Examples 21 and 22.
18	Furthermore, the above polymer can be further diluted by a lower cost oxyger
19	permeable resin, such as EBAC or PE or EVA, down to 50 and even 25% of
20	the original concentration and still maintain a high oxygen scavenging rate, as
21	the non-limiting examples in Examples 23 and 24 show.
00	
22 23	Example 15
24	Polymer Preparation (C1641-6)
	•
25	550 ml of decalin® was placed in a flask. To this was added 350 g of
26	Chevron EMAC SP-2260 which has 24 weight % of methyl acrylate
27	(0.9767 moles of methyl acrylate) and 0.48 g of lrganox®1076 (0.1 mole).
28	The temperature of the mixture was gradually raised while stirring. When the

- 1 temperature reached approximately 120°C, 127.1 g (0.9767 moles) of
- 2 3-methyl-cyclohex-1-ene-4-methanol (97%) was added. When the
- 3 temperature reached approximately 140°C, 4.8 g of the catalyst Ti(OC₂H₅)₄
- 4 was added a portion at a time. The temperature was maintained at 170°C
- 5 while stirring. The course of the reaction was observed by subjecting
- 6 samples of the mixture to NMR at hourly intervals. The percent conversion is
- 7 given in Table 1 below. After 5 hours of reaction, the mixture was cooled and
- 8 400 ml of CHCl₃ was added and the mixture was then precipitated by adding
- 9 it to 4 liters of CH₃OH in a Waring blender. The precipitate is filtered and
- 10 washed with CH₃OH and dried in a vacuum oven at 50°C. The dried mixture
- 11 yielded 407.5 g of ethylene/methyl acrylate/methyl cyclohexene methyl
- 12 acrylate (EMCM).

Table 1

Time (hours)	Percent Conversion 50% 62.3%		
1 hour			
2 hours			
3 hours	65.5%		
5 hours	87.1%		

14

- 15 390 grams of a combination of the above prepared polymer and the same
- 16 polymer prepared under the same conditions in a different batch, which
- 17 together have a conversion percentage of 68.8%, was solvent coated with
- 18 3.25 g cobalt-neodecanoate in 70 ml normal hexane. The mixture was
- 19 tumble dried for 1.5 hours and residual solvent removed in a vacuum.

2021

Example 16

22

Polymer Preparation

- 23 600 ml of decalin was placed in a flask. To this was added 334 grams of
- 24 Chevron SP-2260 (0.9330 moles of methyl acrylate) and 0.44 g of Irganox®

- 1 1010 (0.1% mole). The temperature of the mixture was gradually raised while
- 2 stirring. When the temperature reached approximately 120°C, 104.6 g
- 3 (0.93 moles) of cyclohex-1-ene-4-methanol was added. When the
- 4 temperature reached approximately 140°C, 4.4 g of the catalyst Ti(OC₂H₅)₄
- 5 was added a portion at a time. The temperature was maintained at 160°C
- 6 while stirring. The course of the reaction was observed by subjecting
- 7 samples of the mixture to NMR at hourly intervals. The percent conversion is
- 8 given in Table 2 below. After 3 hours of reaction, the mixture was cooled and
- 9 400 ml of CHCl₃ was added and the mixture was then precipitated by adding
- 10 it to 4 liters of CH₃OH in a Waring blender. The precipitate was filtered and
- 11 washed with CH₃OH and dried in a vacuum oven at 50°C. The dried mixture
- 12 yielded 380.5 g of polymer.

Table 2

Time (hours)	Percent Conversion		
1 hour	43.8%		
2 hours	56.7%		
3 hours	55.7%		

- 15 185 grams of the above-prepared polymer was combined with 45 ml normal
- hexane and 1.54 g cobalt-neodecanoate resulting in 1000 ppm of cobalt ion
- 17 and 0.0185 g Irganox® 1010 resulting in 100 ppm Irganox®. The mixture
- was heated and blended and then dried in a vacuum-oven. The resulting
- 19 compound was extruded into a film.
- 20 Additionally, 185 grams of the above-prepared polymer was combined with
- 21 45 ml normal hexane and 1.54 g cobalt-neodecanoate (resulting in 1000 ppm
- of cobalt ion) and 0.046 g lrganox® 1010 (resulting in 250 ppm lrganox®).
- 23 The mixture is heated and blended and then dried in a vacuum-oven. The
- 24 resulting compound is extruded into a film.

1	Example	17
1	Example	17

2	EMCM Made in ZSK-30 Extruder
3	Ethylene-methyl acrylate copolymer (EMAC) was fed into a Werner &
4	Pfleiderer ZSK-30 twin screw extruder at 6 kg/hr, and the reactants and
5	catalysts were added to the extruder in a subsequent reaction zone. The
6	catalyst Ti(OC ₃ H ₇) ₄ was added with the reactants at 3 mol % or at a rate of
7	148 cc/hr. lrganox®/Toluene solution was added at 4.5 g/900 cc using a
8	Milton Roy 29/290 mini-pump. To obtain 100 ppm of Irganox®, it must be
9	added at 2.2 cc/min. To obtain 50 ppm of Irganox, it must be added at
10	1.1 cc/min. Cyclohexane methyl alcohol with 1,000 ppm of an antioxidant of
11	BHT was added via a Milton Roy dual head at 1958 cc/hr. Steam is injected
12	into the system at 800 cc H₂O/Hr at the end of the reaction zone.
13	51 lbs of EMCM product (100 ppm Irganox® 1010, 59.3% methyl alcohol
14	(MA), 2.98 g/10 min. Melt Flow) was produced over a period of approximately
15	2 hours.

1	Example 18
2	EMCM Made in ZSK-30
3	45 lbs of EMCM product (100 ppm lrganox®1010, 2.38 g/10 min Melt Index)
4	was extruded over a period of approximately 3 hours. A dual steam stripping
5	setup was used in which pressurized injectors at zones 4 and 11 of the
6	extruder pumped steam at 1076 cc/hr and 728 cc/hr, respectively. Both
7	injectors were Pulse 680 pumps with a pressure of at least 800 psi, except at
8	the first measured time interval when injector (No. 4) was measured at
9	500-550 psi and injector (No. 11) was measured at 500 psi.
10	Example 19
11	Co-polymerization of Styrene and 3-Cyclohexene-1-Methanol Methacrylate
12	In a 1-liter round bottom flask, 65 grams styrene (0.625 mole), 113 grams of
13	3-cyclohexene-1-methanol methacrylate (0.625 mole), 1.25 grams of Benzoyl
14	peroxide and 450 grams of toluene were mixed and degassed by freeze-thaw
15	cycles. The degassed solution was polymerized at 70-75°C for 48 hours and
16	discharged into 2 liters of methanol in a Waring Blender. The product isolated
17	was dried in a vacuum oven at 50°C for 2 hours to give 155 grams of
18	co-polymer. NMR analysis indicates it contains 48 mole % of styrene and
19	52 mole % of 3-cyclohexene-1-methanol methacrylate. Tg by DSC is 66°C.
20	
21	Example 20
22	Oxygen Scavenging Test of Styrene/CHMA Copolymer
23	90 weight % of the above-mentioned co-polymer and 10 weight % of a EVA
24	based Master batch containing 1 weight % of co-oleate and 1 weight % of a
25	photoinitiator (BBP³) were processed into a 8 mil thick monolayer film. A
26	100 cm² film was irradiated at both sides to receive 800 m.loutes/cm² of

- 1 254 nm UV on each side and sealed into a foil bag containing 300 cc of 1%
- 2 oxygen. The oxygen uptake was monitored up to 11 days at 4°C and at room
- 3 temperature. The results are shown in Tables 3 (4°C) and 4 (room
- 4 temperature).

Table 3

Time (days)	O₂ Meas. Vol%	O₂ Meas. Vol, ml	Vol-O ₂ Used ml	O₂ Uptake ml/g	O ₂ Uptake Avg Rate cc/m²/day	Instant Rate cc/ m²/day	O ₂ Capacity cc/m²/mil
0.0	1.05	3.15	0.00	0.00	0.00	0.00	0.00
1.1	0.94	2.77	0.32	0.18	14.79	14.79	1.62
3.9	0.49	1.42	1.63	0.92	20.73	23.03	8.15
4.8	0.39	1.11	1.91	1.08	19.81	15.78	9.57
7.0	0.30	0.84	2.17	1.22	15.40	5.72	10.83
11.0	0.09	0.25	2.74	1.54	12.43	7.22	13.72

6

7

Table 4

Time (days)	O₂ Meas. Vol%	O₂ Meas. Vol, ml	Vol-O ₂ Used ml	O₂ Uptake ml/g	O ₂ Uptake Avg Rate cc/m²/day	Instant Rate cc/ m²/day	O ₂ Capacity cc/m²/mil
0.0	1.04	3.12	0.00	0.00	0.00	0.00	0.00
1.1	0.48	1.42	1.65	1.03	75.28	75.28	8.26
3.9	0.09	0.26	2.78	1.73	35.40	19.96	13.92
4.8	0.04	0.11	2.93	1.82	30.26	7.89	14.63
7.0	0.01	0.03	3.01	1.87	21.39	1.91	15.05
11.0	0.01	0.03	3.01	1.87	13.64	0.00	15.05

8

9 Example 21

10

Polymerization of 3-cyclohexene-1-methanol acrylate

WO 99/48963

1	75 grams (0.45 mole) of 3-cyclohexene-1-methanol acrylate (CHAA), 200 ml								
2	of toluene and 0.5 grams of Benzoyl peroxide were charged into a 500 ml								
3	round-bottomed flask and degassed by freeze-thaw cycles. The degassed								
4	solution was polymerized at 70-75°C for 48 hours. The viscous polymer								
5	solution was worked up by precipitating in methanol solution in a Waring								
6	blender. After vacuum drying at room temperature for 3 days, the product is a								
7 °	rubbery clear polymer which weighs 53 grams.								
8									
9	Example 22								
10	Headspace Analysis of O ₂ Scavenging in								
11	Dowlex® 3010/EMCM/Dowlex® 3010 Films								
12	Oxygen scavenging analysis was performed using a Mocon HS750 with a								
13	headspace volume of 300 cc. The sample tested was a 0.48 g three-layer								
14	film with Dowlex® 3010 film for the two outside layers and steam stripped								
15	EMCM (59% converted) for the middle layer (50 ppm Irganox® 1010). The								
16	thickness of the layers was 0.5/1/0.5 +/- 0.1 Mil. The oxygen scavenging								
17	portion of the middle layer comprised 1000 ppm Cobalt salt, 1000 ppm BBP ³								
18	and was exposed for 1.6 minutes to 254 nm UV at 1 inch to receive								
19	800 mJ/cm². The oxygen scavenging was tested with 300 cc 1% O ₂ at 4°C.								
20	The results of the tests are given below in Table 5. These results are plotted								
21	along with the results of Example 20 in Figure 2, which graphically plots %								
22	oxygen in headspace against time (days). The oxygen scavenging uptake								
23	capacity is based on the total weight of the three-layer film.								

-100-

1

Table 5

Time (days)	O ₂ Meas. Vol%	O₂ Meas. Vol, ml	Vol—O ₂ Used ml	O₂ Uptake ml/g	O ₂ Uptake Avg Rate cc/m²/day	Instant Rate cc/ m²/day	O₂ Capacity cc/m²/mil
0.0	1.04	3.12	0.00	0.00	0.00	0.00	0.00
0.8	0.74	2.18	0.89	1.84	57.93	57.93	44.25
1.9	0.46	1.33	1.70	3.54	45.85	37.36	84.85
3.0	0.29	0.83	2.18	4.54	36.87	21.87	109.08
5.8	0.14	0.39	2.60	5.42	22.46	7.41	130.08
7.8	0.09	0.25	2.74	5.71	17.67	3.51	136.95 ·

2

3

15

Example 23

4 Headspace Analysis of O₂ Scavenging in 5 Dowlex® 3010/EMCM/Dowlex® 3010 Films 6 Oxygen scavenging analysis was performed using a Mocon HS750 with a 7 headspace volume of 300 cc. The sample tested was a 0.47 g three-layer 8 film with Dowlex® 3010 film for the two outside layers and steam stripped 9 EMCM for the middle layer (50 ppm Irganox 1010)). The thickness of the layers was 0.5/1/0.5 +/- 0.1 Mil. The oxygen scavenging portion of the middle 10 layer comprised 1000 ppm Cobalt salt, 1000 ppm BBP³ (a photoinitiator) 11 12 exposed for 1.6 minutes to 254 nm UV at 1 inch to receive 800 mJ/cm². The 13 oxygen scavenging was tested with 300 cc 1% O₂ at 4°C. These results are 14 plotted along with the results of Example 21 in Figure 2, which graphically

plots % oxygen in headspace against time (days).

ı	Example 24
2	Headspace Analysis of O ₂ Scavenging in Dowlex® 3010/
3	EBAC:EMCM/Dowlex® 3010 Films
4	Oxygen scavenging analysis was performed using a Mocon HS750 with a
5	headspace volume of 300 cc. The sample tested was a 0.45 g three-layer
6	film with Dowlex® 3010 film for the two outside layers and 3:1 EBAC
7	(ethylene/butyl acrylate copolymer):EMCM (ethylene/methyl acrylate/
8	cyclohexenyl methyl acrylate) for the middle layer (50 ppm Irganox® 1010)).
9	The thickness of the layers was 0.5/1/0.5 +/- 0.1 Mil. The oxygen scavenging
10	portion of the middle layer comprised 1000 ppm of Cobalt salt, 1000 ppm
11	BBP³ was exposed for 1.6 minutes to 254 nm UV at 1 inch to receive
12	800 mJ/cm ² . The oxygen scavenging was tested with 300 cc 1% O ₂ at 4°C.
13	These results are plotted along with the results of Example 22 in Figure 3,
14	which graphically plots % oxygen in headspace against time (days).
15	
16	Example 25
17	Headspace Analysis of O ₂ Scavenging in Dowlex® 3010/
18	EBAC:EMCM/Dowlex® 3010 Films
19	Oxygen scavenging analysis was performed using a Mocon HS750 with a
20	headspace volume of 300 cc. The sample tested was a 0.47 g three-layer
21	film with Dowlex® 3010 film for the two outside layers and 1:1 EBAC:EMCM
22	for the middle layer (50 ppm lrganox®1010)). The thickness of the layers
23	was 0.5/1/0.5 +/- 0.1 Mil. The oxygen scavenging portion of the middle layer
24	comprised 1000 ppm Cobalt Oleate salt, 1000 ppm BBP ³ exposed for
25	1.6 minutes at 254 nm UV at 1 inch to receive 800 mJ/cm². The oxygen
26	scavenging was tested with 300 cc 1% $\rm O_2$ at 4°C. The results of the tests are
27	given below in Table 6. These results are plotted along with the results of
28	Example 23 in Figure 3, which graphically plots % oxygen in headspace

WO 99/48963

-102-

PCT/US99/06379

- 1 against time (days). The oxygen scavenging uptake capacity is based on the
- 2 total weight of the 3-layer film.

Table 6

Time (days)	Head- space O ₂ (Vol%)	Head- space O ₂ (Vol, ml)	Vol—O ₂ Used (ml)	O ₂ Uptake (ml/g)	O₂ Uptake Avg Rate (cc/m²·day)	Instant Rate (cc/m²- day)	O ₂ Capacity cc/m²
0.0	1.09	3.27	0.00	0.00	0.00	0.00	0.00
0.8	0.74	2.18	1.03	2.20	63.00	63.00	51.63
1.8	0.50	1.45	1.73	3.68	48.52	36.18	86.43
4.8	0.17	0.48	2.67	5.8	27.73	15.51	133.45
6.1	0.12	0.34	2.81	5.98	23.17	5.60	140.45
6.9	0.10	0.28	2.86	6.09	20.84	3.40	143.20
7.9	0.08	0.22	2.92	6.21	18.46	2.62	145.90

4 5

6

7

8

9

10

11

12 13

14

15

16

17

18

19

against time (days).

Example 26

Headspace Analysis of O₂ Scavenging Capacity

in Dowlex® 3010/EMCM/Dowlex® 3010 Films

Oxygen scavenging analysis was performed using a Mocon HS750 with a headspace volume of 300 cc. The sample tested was a 0.47 g three-layer film with Dowlex® 3010 film for the two outside layers and steam stripped EMCM for the middle layer (50 ppm Irganox®1010)). The thickness of the layers was 0.5/1/0.5 +/- 0.1 Mil. The oxygen scavenging portion of the middle layer comprised 1000 ppm Cobalt Oleate salt, 1000 ppm BBP³ exposed for 1.6 minutes to 254 nm UV at 1 inch to receive 800 mJ/cm². The oxygen scavenging was tested with 300 cc air at room temperature. The O₂ uptake capacity is based on total weight of the 3-layer film. The results of the tests are given below in Table 7. These results are plotted along with the results of Example 26 in Figure 4, which graphically plots % oxygen in headspace

WO 99/48963

-103-

Table 7

Time (days)	Head- space O ₂ (Vol%)	O ₂ Vol, (ml)	Vol-O₂ Used (ml)	O₂ Uptake (ml/g)	O₂ Uptake Avg Rate (cc/m²·day)	Instant Rate (cc/m²·day)	O ₂ Capacity (cc/m²)
0.0	20.60	61.80	0.00	0.00	0.0	0.0	0.0
1.0	13.40	39.53	21.24	43.35	1058	1058	1062
2.0	12.20	35.38	24.72	50.45	616	173	1236
3.0	11.80	33.63	25.86	52.78	437	60	1293
6.2	11.80	33.04	25.86	52.78	207	0.0	1293

Example 27

1

3

9

10

11 12 Headspace Analysis of O₂ Scavenging Capacity in

4 Dowlex® 3010/EBAC:EMCM/Dowlex® 3010 Films

5 Oxygen scavenging analysis was performed using a Mocon HS750 with a

6 headspace volume of 300 cc. The sample tested was a 0.45 g three-layer

7 film with Dowlex® 3010 film for the two outside layers and 2:1 EBAC:EMCM

8 for the middle layer (50 ppm Irganox 1010)). The width of the layers was

0.5/1/0.5 +/- 0.1 Mil. The oxygen scavenging portion of the middle layer

comprised 1000 ppm of Cobalt salts, 1000 ppm BBP³ exposed for

1.6 minutes to 254 nm UV at 1 inch to receive 800 mJ/cm². The oxygen

scavenging was tested with 300 cc air at room temperature. The O₂ uptake

capacity is based on total weight of the 3-layer film. The results of the tests

14 are given below in Table 8. These results are plotted along with the results of

15 Example 23 in Figure 4, which graphically plots % oxygen in headspace

16 against time (days).

-104-

1 Table 8

Time (days)	Head- space O ₂ (Vol%)	Head- space O₂ Vol, (ml)	Vol-O₂ Used (ml)	O₂ Uptake (ml/g)	O ₂ Uptake Avg Rate (cc/m²-day)	Instant Rate (cc/m²-day)	O ₂ Capacity (cc/m²)
0.0	20.60	61.80	0.00	0.00	0.0	0.0	0.0
1.0	17.70	52.21	8.56	18.20	426	426	428
2.0	17.40	50.46	9.43	20.05	235	43	471
3.0	17.10	48.74	10.28	21.87	174	45	514
6.2	17.10	47.88	10.28	21.87	83	0.0	514

2 3

Example 28

4 Taste Preference Test

5 The organoleptic quality of a film containing EMCM as the scavenging resin in

6 a multi-layer oxygen scavenging packaging structure was evaluated and

7 compared with an SBS (styrene/butadiene/styrene)-based oxygen

8 scavenging packaging structure. Films were triggered with 800 mJ/cm² of

9 254 nm UV. Packages containing ca. 200 ml of water were made and

vacuum/gas flushed to obtain a gas composition of 1% O₂:99% N₂. Packages

were stored at 40°F for seven days prior to taste testing. A forced preference

12 double blind Triangle taste test was carried out on water extracts of the

13 EMCM-based and SBS-based films.

14 15

16

17

18

19

20

21

Sensory results indicated that there was a significant difference (24 out of 28 respondents) between the EMCM-based and SBS-based structures. All 24 respondents who correctly identified the odd sample in the single test preferred the taste of the water packaged in EMCM over SBS. As shown in Table 9, Day 4 scavenging rates of the EMCM-based structures were lower than the SBS counterpart. On Day 4, both structures had significant oxidation and the obvious difference in flavor perception was attributed to the fewer and

- 1 less objectionable by-products (fragments after oxidation of EMCM) of the
- 2 EMCM oxygen scavenging system.
- 3 In a second forced preference triangle taste test, water samples in
- 4 EMCM-based scavenging structures were tested against water samples
- 5 packaged in a standard barrier laminate film (R660B manufactured by
- 6 Cryovac Division of Sealed Air Corporation). The packaged water extract
- 7 samples were submitted to a sensory panel for forced preference double blind
- 8 taste testing. Samples were tested after 8 days of scavenging. A significant
- 9 difference in the taste was found between the samples packaged in the
- 10 EMCM and the control packages. Surprisingly, the preference was towards
- 11 the EMCM structure. Open comments stated that there was no off-flavor
- 12 (normally associated with the SBS-based oxygen scavenging films) in the
- 13 EMCM samples and that EMCM was "pretty close in taste to the control."
- 14 Headspace oxygen levels reached by the EMCM structure were ca. 0.2%
- 15 (down from 1%) at Day 8. Scavenging results of the EMCM film used during
- this test are also listed in Table 9.

Table 9
Oxygen Scavenging
Packaging Films

Film Sample	Average Rate (cc/m²·day)	Average Rate (cc/m²·day)	Induction Period (days)	Peak Instantaneous Rate avg. (c) (cc/m²-day)	Peak Instantaneous Rate (cc/m²-day)
	Mean	St. dev.		Mean	St. dev.
SBS Film	51.0ª	7.8	<1	88.4 (1)	14.1
1 st Sensory test EMCM	41.6°	5.3	<1	68.6 (2)	11.4
2 nd Sensory Test EMCM	30.5 ^b	5.9	<1	83.6 (2-3)	19.4

20

*Rate at 4 days.

21

^bRate at 8 days.

22

Time to reach peak rate in days.

1 Example 29

- 2 Taste Preference Test
- 3 Oxygen scavenging test films, 5 cm x 20 cm, were irradiated with 800 mJ/cm²
- 4 ultraviolet (254 nm) and heat tacked to the top of the test pouches (one per
- 5 pouch). The pouches (16 cm x 19 cm) were made from laminated barrier film
- 6 specifically designed to be oxygen impermeable. 21 gram slices of freshly
- 7 sliced turkey roll were put into sterilized 9 cm petri dishes (one per dish). The
- 8 dishes were, in turn, placed into the barrier pouches (one per pouch). The
- 9 pouches were heat sealed, filled with 300 cc 1% oxygen/99% nitrogen gas,
- 10 and stored at 4°C for the duration of the test.
- 11 Two types of oxygen scavenging polymers were compared in the test against
- 12 a control (barrier pouch alone, no oxygen scavenger). The oxygen
- 13 scavenging films were each three layer (ABA) structures in which the outer,
- 14 "A", layer was 0.5 mil thick LLDPE, and the middle, "B", layer was 1.0 mil thick
- 15 oxygen scavenging polymer (compounded with 1000 ppm cobalt (as oleate)
- 16 and 1000 ppm of a photoinitiator (BBP³). The headspace oxygen for the
- 17 pouches is shown in Table 10. Both of the test oxygen scavenging films
- 18 scavenged more oxygen than the packaged turkey itself.

19 <u>Table 10</u>

Oxygen scavenging layer composition	Initial headspace oxygen, %	Headspace oxygen after 3 days @ 4°C, %
none	1.02	0.72
SBS	1.00	0.08
EMCM	1.02	0.17

20

21 Taste panelists were instructed to force rank the samples according to their

22 taste preference; assigning the least preferred sample a score of 1, and the

- 1 most preferred sample a score of 10. As is shown in Figure 5, the panelists
- 2 found the taste of the turkey packaged in control and the EMCM pouches
- 3 statistically equivalent. The turkey packaged in the SBS pouch was found
- 4 significantly less preferred than either the control or the EMCM.

5 Example 30

- 6 Polymerization of EMCM via high pressure autoclave reactor proceeds in a
- 7 steady-state continuous manner as follows. Ethylene is circulated at a rate of
- 8 10,000-14,000 lb/hr by a hypercompressor which compresses the ethylene to
- 9 16,500-22,500 psig. The compressed ethylene is injected into the autoclave
- 10 reactor in various positions along the reactor wall associated with the zone
- 11 divisions made by the reactor internals. Simultaneously, acrylate of
- 12 cyclohexene-1-methanol (CHAA) comonomer is injected into either the first
- zone or the first and second zones of the reactor at a rate sufficient to
- 14 produce a copolymer containing from 5 to 40% CHAA, more typically
- 15 10%-25% by weight. The reaction is initiated by injection of a solution of
- 16 di-tert butyl peroxypivalate in an aliphatic solvent which also functions as a
- 17 chain transfer agent. The initiator is injected at a rate to provide
- approximately 10-20 ppm (wt) of initiator in the compressed ethylene.
- 19 The locations of the CHAA injection are critical to the polymer being
- 20 produced, as is shown in U.S. Patent No. 5,571,878 which details the effects
- 21 of acrylate injection location on the polymerization of ethylene and an alkyl
- 22 acrylate comonomer in a high pressure system.

- 24 The resultant polymer exits the reactor at a rate of 1000-2000 lb/hr in a
- 25 multi-phase solution in ethylene to a high pressure separator. The pressure
- 26 of the product is reduced adiabatically through a valve to 2,000 psig pressure
- 27 and the unreacted ethylene and unreacted CHAA are recompressed to
- reactor pressure and reinjected into the reactor for further polymerization.
- 29 Additional ethylene is added to the cycle via a primary compressor which

1	compresses the ethylene from pipeline pressure to the suction pressure of the
2	hypercompressor at a rate equal to the polymer production rate.
3	From the high pressure separator, the polymer is reduced in pressure to
4	4-10 psig for further removal of unreacted ethylene and unreacted
5	comonomer. The polymer is fed into a melt pumping device (either an
6	extruder or a gear pump) and is pelletized and transferred for packaging and
7	shipment.
8	
9	Example 31
10	Synthesis of 3-Cyclohexene-1,1-Dimethanol
11	One hundred (100) parts by weight of a formaldehyde aqueous solution
12	(37 wt. % formaldehyde) was charged to a reactor. To this solution, cooled
13	externally with an ice-water bath, was added 118 parts of an aqueous sodium
14	hydroxide solution (25 wt. % sodium hydroxide) by several portions and the
15	temperature of the reaction content was maintained at 20 to 30°C. This was
16	followed by a slow addition of 54 parts of 1,2,5,6-tetrahydrobenzaldehyde at
17	such a rate that the reaction content temperature did not exceed 55°C. After
18	the exotherm dissipated, it was heated at 55°C for two hours with an external
19	heating. The product precipitated out of the solution upon cooling and was
20	collected by suction filtration. The wet-cake was washed thoroughly in the
21	funnel with copious amount of water (5 X 100 parts). The crude product was
22	allowed to dry in air overnight and purified by a recrystallization from toluene.
23	The final product was an off-white colored crystalline material (yield 70%.
24	m.p.: 92-93°C).
25	
26	Example 32
27	Synthesis of 4-Cyclohexene-1,2-Dimethanol
28	A solution of one hundred (100) parts by weight of a 1,2,3,6-
29	tetrahydrophthalic anhydride in 500 parts of dry tetrahydrofuran is slowly
30	added to a stirring mixture of 28.75 parts of lithium aluminum hydride and

1	162 parts of tetranydrofuran. After the addition is complete, the mixture is
2	refluxed for 24 hours. It is then hydrolyzed by a slow addition of a saturated
3	Rochelle salt solution until it turns white. The mixture is refluxed for an
4	additional 10 hours, allowed to room temperature, and suction filtered. The
5	solvent is removed by a distillation and the viscous liquid crude product is
6	purified by a fractional distillation under vacuum (yield 82%. b.p.: 165-170°C
7	at 12 mm).
8	
9	Example 33
10	Synthesis of trans-Diethyl 1,2,3,6-Tetrahydrophthalate
11	One hundred (100) parts by weight of a butadiene is dissolved into a solution
12	of 153 parts of diethyl fumarate in 650 parts of benzene at 0°C. The reaction
13	solution is then heated in a bomb at 50°C for 24 hours. The solvent is
14	removed by a distillation and the liquid crude product is purified by a fractional
15	distillation under vacuum (b.p.: 102-105°C at 2 mm).
16	
17	Example 34
18	Synthesis Of Polyester Containing 3-Cyclohexene-1,1-Dimethanol
19	Dimethyl terephthalate (81.9 g), ethylene glycol (43.7 g), 3-cyclohexene-
20	1,1-dimethanol (20.0 g), and titanium butoxide (0.15 g) were charged into a
21	250 mL 4-necked flask equipped with a distillation column/partial condenser.
22	The agitator and heat were turned on under nitrogen sparge (5 ml/min).
23	When the temperature reached 140-170°C, the methanol collection was
24	started. The temperature was slowly increased to 230°C. The reaction
25	temperature was held at 230-240°C until greater than 95% of the methanol
26	was collected during the course of 2-3 hours at 250-260°C under a full
27	vacuum (0.5-2 mm Hg). The final polyester was discharged into an aluminum
28	pan at about 200°C under nitrogen protection. NMR showed that the
29	polyester contained about 22 wt. % 3-cyclohexene-1,1-dimethanol unit. DSC

1 showed that the polyester was totally amorphous and had a glass transition 2 temperature of 82°C. 3 4 Example 35 5 Synthesis Of Polyester Containing 3-Cyclohexene-1,1-Dimethanol 6 Dimethyl terephthalate (1165.2 g), ethylene glycol (621.0 g), 3-cyclohexene-7 1,1-dimethanol (284.4 g), zinc acetate dihydrate (2.08 g), and antimony oxide 8 (0.62 g) are charged into a 3-liter reaction kettle equipped with a distillation 9 column/partial condenser. The agitator and heat are turned on under 10 nitrogen sparge (10-30 ml/min). When the temperature reaches 140-170°C. 11 the methanol collection is started. After 1-3 hours at 160-190°C under 12 nitrogen, the temperature is slowly increased to 230°C. The reaction 13 temperature is held at 230-240°C until greater than 95% of the methanol is 14 collected during the course of 2-6 hours. Triphenyl phosphite (1.0 a) is then added. The temperature is increased to 250-270°C, the nitrogen is stopped 15 and vacuum is applied. The reaction mixture is held for 2-4 hours at 16 17 250-270°C under a full vacuum (0.5-2 mm Hg). The final polyester is 18 discharged into an aluminum pan at about 200°C under nitrogen protection. 19 20 Example 36 21 Synthesis Of Polyester Containing 3-Cyclohexene-1,1-Dimethanol 22 Following the procedure described in Example 35, dimethyl terephthalate 23 (776.8 g), 1,3-propanediol (304.4 g), 3-cyclohexene-1,1-dimethanol (284.4 g), 24 and titanium butoxide (1.3 g) are charged into a 3-liter reaction kettle 25 equipped with a distillation column/partial condenser. Triphenyl phosphite 26 (0.8 g) is added before increasing the reaction temperature from 230-240°C 27 to over 250°C and applying vacuum.

1	Example 37
2	Synthesis of Polyester Containing 1,2,3,6-Tetrahydrophthalic Acid
3	Ethylene glycol (248.1 g), 1,2,3,6-tetrahydrophthalic anhydride (456.6 g),
4	hydrated monobutyltin oxide (0.7 g), and triphenyl phosphite (0.35 g) were
5	charged into a 2-liter reaction flask equipped with a distillation column/partial
6	condenser. The agitator and heat are turned on under nitrogen sparge
7	(10-30 ml/min). When the temperature reaches 160-180°C, the water
8	collection was started. After 1-3 hours at 160-190°C under nitrogen, the
9	temperature was slowly increased to 230°C. The reaction temperature was
10	held at 230-240°C until greater than 95% of the water was collected during
11	the course of 2-6 hours. The temperature was increased to 250-270°C, the
12	nitrogen was stopped and vacuum was applied. The reaction mixture was
13	held for 2-4 hours at 250-270°C under a full vacuum (0.5-2 mm Hg). The fina
14	polyester was discharged into an aluminum pan at about 200°C under
15	nitrogen protection. NMR confirmed that the polyester was a
16	tetrahydrophthalic acid/ethylene glycol homopolyester. DSC showed that the
17	polyester was totally amorphous and had a glass transition temperature of
18	27°C.
19	Example 38
20	Synthesis Of Polyester Containing 3-Cyclohexene-1,1-Dimethanol
21	and 1,2,3,6-Tetrahydrophthalic Acid
22	Following the procedure described in Example 37, ethylene glycol (248.4 g),
23	1,2,3,6-tetrahydrophthalic anhydride (913.2 g), 3-cyclohexene-1,1-dimethano
24	(839.0 g), and hydrated monobutyltin oxide (1.0 g) are charged into a 3-liter
25	reaction kettle equipped with a distillation column/partial condenser.
26	Triphenyl phosphite (1.0 g) is added before increasing the reaction
27	temperature from 230-240°C to over 250°C and applying vacuum.

1	Example 39
2	Synthesis Of Polyester Containing 3-Cyclohexene-1,1-Dimethanol
3	and 1,2,3,6-Tetrahydrophthalic Acid
4	Following the procedure described in Example 37, 2-methyl-1,3-propanediol
5	(360.4 g), 1,2,3,6-tetrahydrophthalic anhydride (913.2 g), 3-cyclohexene-
6	1,1-dimethanol (839.0 g), and hydrated monobutyltin oxide (1.0 g) are
7	charged into a 3-liter reaction kettle equipped with a distillation column/partial
8	condenser. Triphenyl phosphite (1.0 g) is added before increasing the
9	reaction temperature from 230-240°C to over 250°C and applying vacuum.
10	
11	Example 40
12	Synthesis of Polyester Containing 1,2,3,6-Tetrahydrophthalic Acid
13	Following the procedure described in Example 37, 2 methyl-1,3-propanediol
14	(720.8 g), 1,2,3,6-tetrahydrophthalic anhydride (913.2 g), and hydrated
15	monobutyltin oxide (0.82 g) are charged into a 3-liter reaction kettle equipped
16	with a distillation column/partial condenser. Triphenyl phosphite (0.82 g) is
17	added before increasing the reaction temperature from 230-240°C to over
18	250°C and applying vacuum.
19	
20	Example 41
21	Synthesis of Polyester Containing 1,2,3,6-Tetrahydrophthalic Acid
22	Following the procedure described in Example 37, 1,3-propanediol (608.8 g),
23	1,2,3,6-tetrahydrophthalic anhydride (913.2 g), and hydrated monobutyltin
24	oxide (0.76 g) are charged into a 3-liter reaction kettle equipped with a
25	distillation column/partial condenser. Triphenyl phosphite (0.76 g) is added
26	before increasing the reaction temperature from 230-240°C to over 250°C
27	and applying vacuum.
28	
29	
30	

1 Example 42 2 Synthesis Of Polyester Containing 3-Cyclohexene-1,1-Dimethanol 3 Following the procedure described in Example 37, 2-methyl-1,3-propanediol 4 (180.2 g), adipic acid (584.4 g), 3-cyclohexene-1,1-dimethanol (569.6 g), and 5 hydrated monobutyltin oxide (0.67 g) are charged into a 3-liter reaction kettle 6 equipped with a distillation column/partial condenser. Triphenyl phosphite 7 (0.67 g) is added before increasing the reaction temperature from 230-240°C 8 to over 250°C and applying vacuum. 9 10 Example 43 11 Synthesis Of Polyester Containing 3-Cyclohexene-1,1-Dimethanol 12 Terephthalic acid (664.4 g), 3-cyclohexene-1,1-dimethanol (284.8 g), 13 2-methyl-1,3-propandiol (360.4 g), and hydrated monobutyltin oxide (0.75 g) 14 are charged into a 3-liter reaction kettle equipped with a distillation 15 column/partial condenser. The agitator and heat are turned on under 16 nitrogen sparge (10-30 ml/min). When the temperature reaches 200-220°C, 17 the water collection is started. After 3-7 hours at 200-230°C under nitrogen, 18 the temperature is increased to 240°C. The reaction temperature is held at 19 240°C until greater than 95% of the water is collected during the course of 20 2-6 hours. Triphenyl phosphite (0.75 g) is then added. The temperature is 21 increased to 250-270°C, the nitrogen is stopped and vacuum is applied. The 22 reaction mixture is held for 2-4 hours at 250-270°C under a full vacuum 23 (0.5-2 mm Hg). The final polyester is discharged into an aluminum pan at 24 about 200°C under nitrogen protection. 25 26 Example 44 27 Polymer prepared in Example 34 was solvent cast into a 3.5 mil film 28 containing 2 wt. % cobalt in the form of cobalt cleate and 2 wt. % of 29 anthraquinone as a long wavelength photoinitiator. A 5 x 20 cm² size film was

1	cut and irradiated under a 450 watts medium pressure mercury UV lamp for
2	2 minutes prior to sealing into a foil pouch filled with 300 cc of 1% oxygen.
3	The headspace analysis after 1 day at room temperature showed a reduction
4	in oxygen concentration to 0.91%.
5	
6	Example 45
7	Polymer prepared in Example 37 was solvent cast into 2 mil film containing
8	0.2 wt.% cobalt in the form of cobalt oleate and 1 wt.% of anthraquinone.
9	Irradiated under a 450 watts medium pressure mercury UV lamp for 2 minutes
10	prior to sealing into a foil pouch filled with 300 cc of 1% oxygen. The
11	headspace analysis after 4 days at room temperature showed a reduction in
12	oxygen concentration to 0.83%.
13	
14	Also included within this example and the scope of the invention are
15	compositions comprising various combinations of these substances and
16	materials.
17	
18	Aspects of the present invention have been described by way of example only
19	and it should be appreciated that modifications and additions may be made
20	thereto without departing from the scope thereof.

1 WHAT IS CLAIMED:

2

4

5

6 7

8 9 10

- 1. A composition suitable for scavenging oxygen comprising a mixture of:
 - (a) a polymer or lower molecular weight material containing substituted cyclohexene functionality according to the following structure (I):

 $(I) \qquad \begin{array}{c} A \\ B \\ B \end{array}$

11

where A may be hydrogen or methyl and either one or two of the B groups is a heteroatom containing linkage which attaches the cyclohexene ring to the said material. The remaining B groups are hydrogen or methyl.

16

(b) a transition metal catalyst.

17 18

19 2. The composition of claim 1 where the material is blended with a carrier20 resin.

21

- 22 3. The composition according to claim 2 wherein said mixture further23 contains at least one photoinitiator.
- The composition according to claim 1 wherein the heteroatom
 containing linkage contains an ester, ether, amide, imide, urethane, or acetal
 group.

27

5. An oxygen scavenger composition comprising a polymer or oligomer
having at least one cyclohexene group, and a transition metal salt, compound
or complex.

- -116-
- 1 6. The composition of claim 5 further comprising a trigger enhancing
- 2 component which makes the scavenger susceptible to triggering from an
- 3 external even.
- 4 7. The composition of claim 6, wherein the trigger enhancing component is
- 5 selected from the group consisting of benzophenone or substituted
- 6 benzophenone.
- 7 8. The composition of claim 6, wherein the external event is irradiation by
- 8 electromagnetic radiation.
- 9 9. The composition of claim 6, wherein the external event is irradiation by
- 10 UV light.
- 11 10. The composition of claim 5, wherein the material is blended with a
- 12 carrier resin.
- 13 11. The composition of claim 5, wherein the oxygen scavenger composition
- 14 is in the form of a plastics resin.
- 15 12. The composition of claim 11, wherein the plastics resin comprises a
- 16 polyester resin.
- 17 13. The composition of claim 11, wherein the plastics resin comprises a
- resin suitable for use in the manufacture of plastic films.
- 19 14. The composition of claim 5 wherein the composition is prepared from
- 20 the reaction of a tetrahydrophthalic anhydride with at least one of:
- 21 i) a diol;
- 22 ii) a hydroxy compound; or
- 23 iii) a polyhydroxy compound.
- 24 15. The composition of claim 14 wherein the composition is prepared by
- 25 heating tetrahydrophthalic anhydride with at least one of:
- 26 i) a diol:
- 27 ii) a hydroxy compound; or
- 28 iii) a polyhydroxy compound,
- 29 in a solvent.

- 1 16. The oxygen scavenger composition of claim 14 wherein the anhydride
- 2 comprises 1,2,3,6 tetrahydrophthalic anhydride, or tetrahydrophthalic
- 3 anhydride monomer derivable from butadiene.
- 4 17. The composition of claim 14 wherein the composition is prepared by a
- 5 reactive extrusion process.
- 6 18. The composition of claim 5 prepared from the reaction of a
- 7 tetrahydrobenzyl alcohol with one or more compounds having one or more of
- 8 the following functionalities:
- 9 i) carboxylic acid;
- 10 ii) acid halide;
- 11 iii) ester;
- 12 iv) anhydride; and
- 13 v) isocyanate.
- 14 19. The oxygen scavenger composition of claim 18 wherein the composition
- 15 is prepared by a reactive extrusion process.
- 16 20. The oxygen scavenger composition of claim 18 wherein the composition
- 17 is prepared from the reaction of a tetrahydrobenzyl alcohol with an ester by a
- 18 transesterification process.
- 19 21. The oxygen scavenger composition of claim 18 wherein the compound
- 20 with the anhydride functionality comprises styrene maleic anhydride
- 21 copolymer.
- 22 22. The oxygen scavenger composition of claim 18 wherein the compound
- 23 with the isocyanate functionality comprises polyfunctional isocyanate.
- 24 23. The oxygen scavenger composition of claim 5 comprising a polyester,
- wherein the composition is prepared from cyclohexene dimethanol.
- 26 24. The oxygen scavenger composition of claim 5, wherein the composition
- 27 is prepared from tetrahydrobenzoic acid and a hydroxyl functional material.
- 28 25. The oxygen scavenger composition of claim 5, wherein the composition
- 29 is prepared from tetrahydrobenzoic acid and a hydroxyl functional material.

- 1 26. The oxygen scavenger composition of claim 5, wherein the composition
- 2 is prepared from tetrahydrobenzaldehyde and a hydroxyl functional material.
- 3 27. The oxygen scavenger composition of claim 5 comprising a polymer or
- 4 oligomer having at least one cyclohexene group, wherein some carbons of
- 5 the cyclohexene group form part of other ring structures within the polymer or
- 6 oligomer.
- 7 28. The oxygen scavenger composition of claim 5 comprising a pendant
- 8 cyclic alkene group, the composition prepared by a method wherein some
- 9 carbons of the cyclohexene group form part of the skeleton of the polymer or
- 10 oligomer.
- 11 29. The oxygen scavenger composition of claim 5, comprising a pendant
- 12 cyclohexene group, wherein the composition is prepared by a method
- including a Diels Alder addition reaction.
- 14 30. The oxygen scavenger composition of claim 5 wherein the composition
- 15 is incorporated in a sachet.

1 2 31. A composition comprising a polymeric backbone, cyclic olefinic pendent

3 groups and linking groups linking the olefinic pendent groups to the polymeric

4 backbone.

5 32. A composition according to claim 31, wherein the polymeric backbone is

6 ethylenic and the linking groups are selected from the group consisting

7 of:

- 10 wherein R is hydrogen or an alkyl group selected from the group
- 11 consisting of methyl, ethyl, propyl and butyl groups and where n is an
- integer in the range from 1 to 12. 12

33. The composition of claim 31, wherein the cyclic olefinic pendent groups 13

14 have the structure (II):

16

17

(II)
$$q_1 \xrightarrow{q_2} m$$

$$q_4 \xrightarrow{q_3}$$

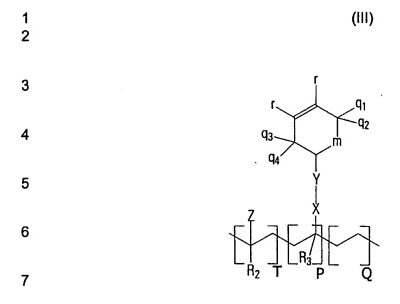
18 where q₁, q₂, q₃, q₄, and r are selected from the group consisting of –H, -

CH₃, and -C₂H₅; and where m is -(CH₂)_n- with n being an integer in the 19-

20 range from 0 to 4; and wherein, when r is -H, at least one of q₁, q₂, q₃

21 and q₄ is –H.

1	38.	The composition of claim 31, wherein the polymeric backbone	
2		comprises monomers selected from the group consisting of ethylene	
3		and styrene.	
4			
5	38.	The composition of claim 31, wherein the cyclic olefinic pendent groups	
6		are grafted onto the linking groups of the polymeric backbone by an	
7		esterification, transesterification, amidation or transamidation reaction.	
8	38.	The composition of claim 35, wherein the esterification,	
9		transesterification, amidation or transamidation reaction is a solution	
10		reaction or a reactive extrusion.	
11	38.	The composition of claim 35, wherein the esterification,	
12		transesterification, amidation or transamidation reaction is catalyzed by	
13		a catalyst selected from the group consisting of strong non-oxidizing	
14		acids, tertiary amines, Group I alkoxides, Group IVB alkoxides, and	
15		Group IVA organometallics.	
16	38.	The composition of claim 37, wherein the catalyst is selected from a	
17		group consisting of toluene sulfonic acid, sodium methoxide, tetrabutyl	
18		titanate, tetraisopropyl titanate, tetra-n-propyl-titanate, tetraethyl titanate,	
19		2-hydroxy-pyridine and dibutyltin dilaurate.	
20	39.	The composition of claim 31, wherein the polymeric backbone, linking	
21	groups and cyclic olefin pendent groups comprise repeating units, each unit		
22	havi	ng a structure (III) as follows:	



wherein P+T+ Q is 100 mol % of the total composition; P is greater than 0
mol % of the total composition; Z is selected from the group consisting
of an aryl group; -(C=O)OR₁; -O(C=O)R₁; and an alkyl aryl group,
structure(IV):

14

15 where R₄ is selected from the group consisting of –CH₃, -C₂H₅, and –H; R₁ is 16 selected from the group consisting of -H, -CH₃, -C₂H₅, -C₃H₇ and -C₄H₉; 17 R₂ and R₃ are selected from the group consisting of -H and -CH₃; X is selected from the group consisting of -O-, -NH-, -(C=O)O-, -(C=O)NH-, 18 -(C=O)S-, -O(C=O)- and -(CHR) $_{\ell}$ -; ℓ is an integer in the range from 1 to 19 20 6; Y is –(CHR)_n-, where n is an integer in the range from 0 to 12, R 21 being selected from the group consisting of -H, -CH₃ and -C₂H₅; where 22 q₁, q₂, q₃, q₄, and r are selected from the group consisting of –H, -CH₃,

- and $-C_2H_5$; and where m is $-(CH_2)_{n-}$ and where n is an integer in the
- 2 range from 0 to 4; and wherein when r is -H, at least one of q₁, q₂, q₃
- 3 and q_4 is -H.
- 4 40. The composition of claim 39, wherein the cyclic olefinic pendent groups
- 5 are selected from the group consisting of cyclohexene-4-methylene radical,
- 6 1-methyl cyclohexene-4-methylene radical, 2-methyl cyclohexene-4-
- 7 methylene radical, 5-methyl cyclohexene-4-methylene radical, 1,2-dimethyl
- 8 cyclohexene-4-methylene radical, 1,5-dimethyl cyclohexene-4-methylene
- 9 radical, 2,5-dimethyl cyclohexene-4-methylene radical, 1,2,5-trimethyl
- 10 cyclohexene-4-methylene radical, cyclohexene-4-ethylene radical, 1-methyl
- 11 cyclohexene-4-ethylene radical, 2-methyl cyclohexene-4-ethylene radical.
- 12 5-methyl cyclohexene-4-ethylene radical, 1,2-dimethyl cyclohexene-4-
- 13 ethylene radical, 1,5-dimethyl cyclohexene-4-ethylene radical, 2,5-dimethyl
- 14 cyclohexene-4-ethylene radical, 1,2,5-trimethyl cyclohexene-4-ethylene
- 15 radical, cyclohexene-4-propylene radical, 1-methyl cyclohexene-4-propylene
- 16 radical, 2-methyl cyclohexene-4-propylene radical, 5-methyl cyclohexene-4-
- 17 propylene radical, 1,2-dimethyl cyclohexene-4-propylene radical, 1,5-dimethyl
- 18 cyclohexene-4-propylene radical, 2,5-dimethyl cyclohexene-4-propylene
- 19 radical, 1,2,5-trimethyl cyclohexene-4-propylene radical, cyclopentene-4-
- 20 methylene radical, 1-methyl cyclopentene-4-methylene radical, 3-methyl
- 21 cyclopentene-4-methylene radical, 1,2-dimethyl cyclopentene-4-methylene
- radical, 3,5-dimethyl cyclopentene-4-methylene radical, 1,3-dimethyl
- 23 cyclopentene-4-methylene radical, 2,3-dimethyl cyclopentene-4-methylene
- radical, 1,2,3-trimethyl cyclopentene-4-methylene radical, 1,2,3,5-tetramethyl
- 25 cyclopentene-4-methylene radical, cyclopentene-4-ethylene radical, 1-methyl
- 26 cyclopentene-4-ethylene radical, 3-methyl cyclopentene-4-ethylene radical,
- 27 1,2-dimethyl cyclopentene-4-ethylene radical, 3,5-dimethyl cyclopentene-4-
- ethylene radical, 1,3-dimethyl cyclopentene-4-ethylene radical, 2,3-dimethyl
- 29 cyclopentene-4-ethylene radical, 1,2,3-trimethyl cyclopentene-4-ethylene

- 1 radical, 1,2,3,5-tetramethyl cyclopentene-4-ethylene radical, cyclopentene-4-
- 2 propylene radical, 1-methyl cyclopentene-4-propylene radical, 3-methyl
- 3 cyclopentene-4-propylene radical, 1,2-dimethyl cyclopentene-4-propylene
- 4 radical, 3,5-dimethyl cyclopentene-4-propylene radical, 1,3-dimethyl
- 5 cyclopentene-4-propylene radical, 2,3-dimethyl cyclopentene-4-propylene
- 6 radical, 1,2,3-trimethyl cyclopentene-4-propylene radical, and 1,2,3,5-
- 7 tetramethyl cyclopentene-4-propylene radical.
- 8 41. The composition of claim 39, wherein the composition is an
- 9 ethylene/methyl acrylate/cyclohexenyl methyl acrylate terpolymer, a
- 10 cyclohexenyl methyl acrylate/ethylene copolymer, a cyclohexenyl methyl
- 11 methacrylate/styrene copolymer, a cyclohexenyl methyl acrylate
- 12 homopolymer or a methyl acrylate/cyclohexenyl methyl acrylate copolymer.
- 13 42. An oxygen scavenging composition comprising a polymeric backbone.
- 14 cyclic olefinic pendent groups, linking groups linking the olefinic pendent
- groups to the polymeric backbone and a transition metal catalyst.
- 16 43. A composition according to claim 42, wherein the polymeric backbone is
- 17 ethylenic and the linking groups are selected from the group consisting
- 18 of:
- 19 -O-(CHR)₀-; -(C=O)-O-(CHR)₀-; -NH-(CHR)₀-; -O-(C=O)-(CHR)₀-;
- 20 -(C=O)-NH-(-CHR)_n-; and -(C=O)-O-CHOH-CH₂-O-;
- 21 wherein R is hydrogen or an alkyl group selected from the group
- consisting of methyl, ethyl, propyl and butyl groups and where n is an
- integer in the range from 1 to 12.
- 24 44. The composition of claim 42, wherein the cyclic olefinic pendent groups
- 25 have the structure (II):

1 (II)

2

3

6

7 where q_1 , q_2 , q_3 , q_4 , and r are selected from the group consisting of -H, -CH₃,

8 and $-C_2H_5$; and where m is $-(CH_2)_n$ - with n being an integer in the range

from 0 to 4; and wherein, when r is -H, at least one of q_1 , q_2 , q_3 and q_4 is

10 -H.

- 11 45. The composition of claim 42, wherein the polymeric backbone
- comprises monomers selected from the group consisting of ethylene
- and styrene.
- 14 46. The composition of claim 42, wherein the cyclic olefinic pendent groups
- are grafted onto the linking groups of the polymeric backbone by a
- esterification, transesterification, amidation or transamidation reaction.
- 17 47. The composition of claim 46, wherein the esterification,
- transesterification, amidation or transamidation reaction is a solution
- 19 reaction or a reactive extrusion.
- 20 48. The composition of claim 46, wherein the esterification,
- transesterification, amidation or transamidation reaction is catalyzed by
- a catalyst selected from the group consisting of strong non-oxidizing

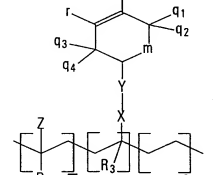
- 1 acids, tertiary amines, Group I alkoxides, Group IVB alkoxides, and
- 2 Group IVA organometallics.
- 3 49. The composition of claim 48, wherein the catalyst is selected from a
- 4 group consisting of toluene sulfonic acid, sodium methoxide, tetrabutyl
- 5 titanate, tetraisopropyl titanate, tetra-n-propyl-titanate, tetraethyl titanate,
- 6 2-hydroxy-pyridine and dibutyltin dilaurate.
- 7 50. The composition of claim 42, wherein the polymeric backbone, linking
- 8 groups and cyclic olefin pendent groups comprise repeating units, each
- 9 unit having a structure (III) as follows:
- 10 (III)

12

13

14

15



- wherein P + T + Q is 100 mol % of the total composition; P is greater than 0
- mol % of the total composition; Z is selected from the group consisting
- of an aryl group; -(C=O)OR₁; -O(C=O)R₁; and an alkyl aryl group,
- 19 structure (IV):

20

(IV)

1 where R₄ is selected from the group consisting of –CH₃, -C₂H₅, and -H; R₁ is 2 selected from the group consisting of -H, -CH₃, -C₂H₅, -C₃H₇ and -C₄H₉; R₂ and R₃ are selected from the group consisting of -H and -CH₃; X is 3 4 selected from the group consisting of -O-, -NH-, -(C=O)O-, -(C=O)NH-, -(C=O)S-, -O(C=O)- and -(CHR),-; ℓ is an integer in the range from 1 to 5 6; Y is -(CHR)₀-, where n is an integer in the range from 0 to 12, R being 6 selected from the group consisting of -H, -CH₃ and -C₂H₅; where q₁, q₂, 7 q₃, q₄, and r are selected from the group consisting of -H, -CH₃, and 8 $-C_2H_{5}$; and where m is $-(CH_2)_0$ - and where n is an integer in the range 9 from 0 to 4; and wherein when r is -H, at least one of q_1 , q_2 , q_3 and q_4 is 10 11 **−**H. 51. The composition of claim 50, wherein the cyclic olefinic pendent groups 12 13 are selected from the group consisting of cyclohexene-4-methylene 14 radical, 1-methyl cyclohexene-4-methylene radical, 2-methyl cyclohexene-4-methylene radical, 5-methyl cyclohexene-4-methylene 15 radical, 1,2-dimethyl cyclohexene-4-methylene radical, 1,5-dimethyl 16 17 cyclohexene-4-methylene radical, 2,5-dimethyl cyclohexene-4methylene radical, 1,2,5-trimethyl cyclohexene-4-methylene radical, 18 19 cyclohexene-4-ethylene radical, 1-methyl cyclohexene-4-ethylene radical, 2-methyl cyclohexene-4-ethylene radical, 5-methyl cyclohexene-20 21 4-ethylene radical, 1,2-dimethyl cyclohexene-4-ethylene radical, 1,5-22 dimethyl cyclohexene-4-ethylene radical, 2,5-dimethyl cyclohexene-4ethylene radical, 1,2,5-trimethyl cyclohexene-4-ethylene radical, 23 cyclohexene-4-propylene radical, 1-methyl cyclohexene-4-propylene 24 25 radical, 2-methyl cyclohexene-4-propylene radical, 5-methyl 26 cyclohexene-4-propylene radical, 1,2-dimethyl cyclohexene-4-propylene 27 radical, 1,5-dimethyl cyclohexene-4-propylene radical, 2,5-dimethyl 28 cyclohexene-4-propylene radical, 1,2,5-trimethyl cyclohexene-4-29 propylene radical, cyclopentene-4-methylene radical, 1-methyl

28

29

1	cyclopentene-4-methylene radical, 3-methyl cyclopentene-4-methylene
2	radical, 1,2-dimethyl cyclopentene-4-methylene radical, 3,5-dimethyl
3	cyclopentene-4-methylene radical, 1,3-dimethyl cyclopentene-4-
4	methylene radical, 2,3-dimethyl cyclopentene-4-methylene radical, 1,2,3-
5	trimethyl cyclopentene-4-methylene radical, 1,2,3,5-tetramethyl
6	cyclopentene-4-methylene radical, cyclopentene-4-ethylene radical, 1-
7	methyl cyclopentene-4-ethylene radical, 3-methyl cyclopentene-4-
8	ethylene radical, 1,2-dimethyl cyclopentene-4-ethylene radical, 3,5-
9	dimethyl cyclopentene-4-ethylene radical, 1,3-dimethyl cyclopentene-4-
10	ethylene radical, 2,3-dimethyl cyclopentene-4-ethylene radical, 1,2,3-
11	trimethyl cyclopentene-4-ethylene radical, 1,2,3,5-tetramethyl
12	cyclopentene-4-ethylene radical, cyclopentene-4-propylene radical, 1-
13	methyl cyclopentene-4-propylene radical, 3-methyl cyclopentene-4-
14	propylene radical, 1,2-dimethyl cyclopentene-4-propylene radical, 3,5-
15	dimethyl cyclopentene-4-propylene radical, 1,3-dimethyl cyclopentene-4-
16	propylene radical, 2,3-dimethyl cyclopentene-4-propylene radical, 1,2,3-
17	trimethyl cyclopentene-4-propylene radical, and 1,2,3,5-tetramethyl
18	cyclopentene-4-propylene radical.
19	52. The composition of claim 50, wherein the composition is an
20	ethylene/methyl acrylate/cyclohexenyl methyl acrylate terpolymer, a
21	cyclohexenyl methyl acrylate/ethylene copolymer, a cyclohexenyl methyl
22	methacrylate/styrene copolymer, a cyclohexenyl methyl acrylate
23	homopolymer or a methyl acrylate/cyclohexenyl methyl acrylate copolymer.
24	53. The composition of claim 42, wherein odor and taste characteristics of
25	products packaged with material comprised of the composition are not
26	adulterated as a result of oxidation of the composition.

54. The composition of claim 42, wherein there is no significant

fragmentation of the olefinic pendent groups and linking groups from the

polymeric backbone as a result of oxidation of the composition.

- 1 54. The composition of claim 42, wherein there is no significant
- 2 fragmentation of the olefinic pendent groups and linking groups from the
- 3 polymeric backbone as a result of oxidation of the composition.
- 4 55. The composition of claim 42, wherein the transition metal catalyst is a
- 5 metal salt.
- 6 56. The composition of claim 55, wherein the metal in the metal salt is
- 7 cobalt.
- 8 57. The composition according to claim 55, wherein the metal salt is
- 9 selected from the group consisting of cobalt neodecanoate, cobalt
- 10 2-ethylhexanoate, cobalt oleate and cobalt stearate.
- 11 58. The composition of claim 42, further comprising at least one triggering
- material to enhance initiation of oxygen scavenging.
- 13 59. The composition of claim 58, wherein the triggering material is a photo
- 14 initiator.
- 15 60. An article of manufacture suitable as a container, the container inhibiting
- oxidation of contents of the container by removing oxygen from the
- 17 container and by inhibiting ingress of oxygen into the container from
- outside the container, wherein the article comprises an oxygen
- scavenging composition which comprises a polymeric backbone, cyclic
- 20 olefinic pendent groups, linking groups linking the olefinic pendent
- groups to the backbone, and a transition metal catalyst.
- 22 61. The article of manufacture of claim 60, wherein the polymeric backbone
- is ethylenic and the linking groups are selected from the group
- 24 consisting of:
- 25 -O-(CHR)_n-; -(C=O)-O-(CHR)_n-; -NH-(CHR)_n-; -O-(C=O)-(CHR)_n-;
- 26 -(C=O)-NH-(-CHR)_n-; and -(C=O)-O-CHOH-CH₂-O-;

- 1 wherein R is hydrogen or an alkyl group selected from the group
- consisting of methyl, ethyl, propyl and butyl groups and where n is an
- 3 integer in the range from 1 to 12.
- 4 62. The article of manufacture of claim 60, wherein the cyclic olefinic
- 5 pendent groups have the structure (II):
- 6 (II)

- where q_1 , q_2 , q_3 , q_4 , and r are selected from the group consisting of -H, -CH₃,
- and $-C_2H_5$; and where m is $-(CH_2)_n$ and where n is an integer in the
- range of from 0 to 4; and wherein when r is -H, at least one of q₁, q₂, q₃
- 13 and q_4 is also -H.
- 14 63. The article of manufacture of claim 60, wherein the polymeric backbone
- 15 comprises monomers selected from the group consisting of ethylene and
- 16 styrene.
- 17 64. The article of manufacture of claim 60, wherein the cyclic olefinic
- pendent groups are grafted onto the linking groups of the polymeric
- backbone by a esterification, transesterification, amidation or
- 20 transamidation reaction.
- 21 65. The article of manufacture of claim 64, wherein the esterification,
- transesterification, amidation or transamidation reaction is a solution
- reaction or a reactive extrusion.

- The article of manufacture of claim 64, wherein the esterification, transesterification, amidation or transamidation reaction is catalyzed by a catalyst selected from the group consisting of strong non-oxidizing acids, tertiary amines, Group I alkoxides, Group IVB alkoxides, and Group IVA organometallics.
- 6 67. The article of manufacture of claim 66, wherein the catalyst is selected from the group consisting of toluene sulfonic acid, sodium methoxide, tetrabutyl titanate, tetraisopropyl titanate, tetra-n-propyl-titanate, tetraethyl titanate, 2-hydroxy-pyridine and dibutyltin dilaurate.
- 10 68. The article of manufacture of claim 60, wherein the backbone, linking
 11 groups and cyclic olefin pendent groups comprise repeating units, each
 12 unit having a structure (III) as follows:

13 (III)

18

23

14
$$q_3 \qquad q_1 \qquad q_2 \qquad q_3 \qquad q_4 \qquad q_5 \qquad q_6 \qquad q$$

wherein P+T + Q is 100 mol % of the total composition; P is greater than 0 mol % of the total composition; Z is selected from the group consisting of an aryl group, -(C=O)OR₁, -O(C=O)R₁ and an alkyl aryl group, structure (IV):

2

3

4

5

14

15

16

17

18

19 20

21

22

23

24

2526

(IV)

V) (1)

6 selected from the group consisting of –H, -CH₃, -C₂H₅, -C₃H₇ and –C₄H₉; 7 R₂ and R₃ are selected from the group consisting of –H and CH₃; X is selected from the group consisting of O-, -NH-, -(C=O)O-, -(C=O)NH-, 8 -(C=O)S-, -O(C=O)- and -(CHR),-; ℓ is an integer selected from the 9 group consisting of 1, 2, 3, 4, 5 and 6; Y is -(CHR), where n is an 10 11 integer in the range from 0 to 12 and R is selected from the group 12 consisting of $-H_1$, $-CH_3$ and C_2H_5 ; and where q_1 , q_2 , q_3 , q_4 , and r are 13 selected from the group consisting of -H, -CH₃, and -C₂H₅; and where m

is $-(CH_2)_n$ - and where n is an integer in the range of from 0 to 4; and

wherein when r is -H, at least one of q_1 , q_2 , q_3 , and q_4 is -H.

where R₄ is selected from the group consisting of --CH₃, -C₂H₅, and --H; R₁ is

69. The article of manufacture of claim 60, wherein the cyclic olefinic pendent groups are selected from the group consisting of cyclohexene-4-methylene radical, 1-methyl cyclohexene-4-methylene radical, 2-methyl cyclohexene-4-methylene radical, 5-methyl cyclohexene-4-methylene radical, 1,2-dimethyl cyclohexene-4-methylene radical, 1,5-dimethyl cyclohexene-4-methylene radical, 1,2,5-trimethyl cyclohexene-4-methylene radical, cyclohexene-4-ethylene radical, 1-methyl cyclohexene-4-ethylene radical, 2-methyl cyclohexene-4-ethylene radical, 5-methyl cyclohexene-4-ethylene radical, 1,2-dimethyl cyclohexene-4-ethylene radical, 1,5-dimethyl cyclohexene-4-ethylene radical, 1,5-dimethyl cyclohexene-4-ethylene radical, 2,5-dimethyl

cyclohexene-4-ethylene radical, 1,2,5-trimethyl cyclohexene-4-ethylene

- 1 radical, cyclohexene-4-propylene radical, 1-methyl cyclohexene-4-propylene 2 radical, 2-methyl cyclohexene-4-propylene radical, 5-methyl cyclohexene-4-3 propylene radical, 1,2-dimethyl cyclohexene-4-propylene radical, 1,5-dimethyl 4 cyclohexene-4-propylene radical, 2,5-dimethyl cyclohexene-4-propylene radical, 1,2,5-trimethyl cyclohexene-4-propylene radical, cyclopentene-4-5 methylene radical, 1-methyl cyclopentene-4-methylene radical, 3-methyl 6 cvclopentene-4-methylene radical, 1,2-dimethyl cyclopentene-4-methylene 7 8 radical, 3,5-dimethyl cyclopentene-4-methylene radical, 1,3-dimethyl 9 cyclopentene-4-methylene radical, 2,3-dimethyl cyclopentene-4-methylene 10 radical, 1,2,3-trimethyl cyclopentene-4-methylene radical, 1,2,3,5-tetramethyl 11 cyclopentene-4-methylene radical, cyclopentene-4-ethylene radical, 1-methyl 12 cyclopentene-4-ethylene radical, 3-methyl cyclopentene-4-ethylene radical. 13 1,2-dimethyl cyclopentene-4-ethylene radical, 3,5-dimethyl cyclopentene-4-14 ethylene radical, 1,3-dimethyl cyclopentene-4-ethylene radical, 2,3-dimethyl 15 cyclopentene-4-ethylene radical, 1,2,3-trimethyl cyclopentene-4-ethylene 16 radical, 1,2,3,5-tetramethyl cyclopentene-4-ethylene radical, cyclopentene-4-17 propylene radical, 1-methyl cyclopentene-4-propylene radical, 3-methyl 18 cyclopentene-4-propylene radical, 1,2-dimethyl cyclopentene-4-propylene 19 radical, 3,5-dimethyl cyclopentene-4-propylene radical, 1,3-dimethyl 20 cyclopentene-4-propylene radical, 2,3-dimethyl cyclopentene-4-propylene
- 70. The article of manufacture of claim 60, wherein the composition is an
 ethylene/methyl acrylate/cyclohexenyl methyl acrylate terpolymer, a
 cyclohexenyl methyl acrylate/ethylene copolymer, a cyclohexenyl methyl
 methacrylate/styrene copolymer, a cyclohexenyl methyl acrylate
 homopolymer or a methyl acrylate/cyclohexenyl methyl acrylate copolymer.

radical, 1,2,3-trimethyl cyclopentene-4-propylene radical, and 1,2,3,5-

tetramethyl cyclopentene-4-propylene radical.

21

22

1 71. The article of manufacture according to claim 60, wherein the transition metal catalyst is a metal salt. 2 3 4 72. The article of manufacture according to claim 71, wherein the metal in 5 6 the metal salt is cobalt. 7 8 73. The article of manufacture of according to claim 71, wherein the metal 9 salt is selected from the group consisting of cobalt neodecanoate, cobalt 10 2-ethylhexanoate, cobalt oleate and cobalt stearate. 11 12 13 74. The article of manufacture of claim 60, further comprising at least one 14 triggering material to enhance initiation of oxygen scavenging. 15 16 75. The article of manufacture of claim 74, wherein the triggering material is 17 a photoinitiator. 18 19 20 76. The article of manufacture of claim 60, wherein odor and taste 21 characteristics of products packaged with material comprised of the 22 composition are not adulterated as a result of oxidation of the composition. 23 24 77. The article of manufacture of claim 60, wherein there is no significant fragmentation of the olefinic pendent groups and linking groups from the 25 polymeric backbone as a result of oxidation of the composition. 26 27 28 78. The article of manufacture of claim 60 wherein the article is a 29 package.

79.

1

The article of manufacture of claim 78, wherein the package

2	comprises a flexible film having a thickness of at most 10 mil or a flexible		
3	sheet having a thickness of at least 10 mil.		
4			
5	80.	The article of manufacture of claim 78, wherein the oxygen	
6	scaven	ging system of the package comprises at least one additional layer	
7	selecte	d from among oxygen barrier layers, polymeric selective layers, and	
8	heat se	al layers.	
9			
10	81.	The article of manufacture of claim 78, wherein the article is a	
11	package with a food product located within the package.		
12	82.	The article of manufacture of claim 78, wherein the article is a	
13	packag	e for packaging a cosmetic, chemical, electronic device, pesticide or a	
14	pharmaceutical composition.		
15			
16	83.	A multi-layer film comprising the article of manufacture according to	
17	claim 6	0, and at least one additional functional layer.	
18			
19	84.	The multi-layer film according to claim 83, wherein at least one	
20	addition	nal layer is selected from among oxygen barrier layers, polymeric	
21	selectiv	e barrier layers, structural layers and heat seal layers.	
22			
23	85.	The multi-layer film according to claim 83, wherein the at least one	
24	additional layer is an oxygen barrier layer.		
25			
26	86.	The multi-layer film according to claim 85, further comprising at least	
27	one po	ymeric selective barrier layer.	
28			
29	87.	The multi-layer film according to claim 85, further comprising at least	
30	one heat seal layer.		

1			
2	88.	The multi-layer film according to claim 85, further comprising at least	
3	one st	ructural layer.	
4		•	
5	89.	The article of claim 60, wherein the article is a rigid container, sealing	
6	gasket	, patch, container closure device, bottle cap, bottle cap insert or	
7	molded	d or thermoformed shape.	
8			
9	90.	The article of claim 89, wherein the molded or thermoformed shape is	
10	a bottle	e or tray.	
11			
12	91.	A layer suitable for scavenging oxygen comprising:	
13	i)	a polymer backbone;	
14	ii) cyclic olefinic pendent groups;	
15	iii) linking groups linking the backbone with the pendent groups; and		
16	iv) a transition metal catalyst.	
17			
18	92.	The layer of claim 91, wherein odor and taste characteristics of	
19	produc	ts packaged with material comprised of the layer are not adulterated as	
20	a resul	t of oxidation of the layer.	
21			
22	93.	The layer of claim 91, wherein there is no significant fragmentation of	
23	the ole	finic pendent groups and linking groups from the polymeric backbone	
24	а	s a result of oxidation of the layer.	
25			
26	94.	A layer according to claim 91, wherein the transition metal catalyst is	
27	a meta	l salt.	
28			
29	95.	A layer according to claim 94, wherein the transition metal in the	
30	metal s	ealt is cobalt.	

1			
2	96.	A layer according to claim 94, wherein the metal salt selected from	
3	the gro	up consisting of cobalt neodecanoate, cobalt 2-ethylhexanoate, cobalt	
4	oleate and cobalt stearate.		
5			
6	97.	A layer according to claim 91, wherein said layer in addition	
7	compri	ses polymeric diluent.	
8			
9	98.	A layer according to claim 97, wherein said diluent is a thermoplastic	
10	polyme	er.	
11	99.	A layer according to claim 91, wherein said layer is adjacent to one or	
12	more a	dditional layers.	
13			
14	100.	A layer according to claim 99, wherein at least one additional layer is	
15	an oxy	gen barrier.	
16			
17	101.	A layer according to claim 100, wherein said oxygen barrier	
18	comprises a member of the group consisting of poly(ethylene-vinyl alcohol),		
19	polyacrylonitrile, poly(vinyl chloride), polyamides, poly(vinylidene dichloride),		
20	poly(ethylene terephthalate), silica, metal foil and metalized polymeric films.		
21			
22	102.	A layer according to claim 99, wherein one or more of said additional	
23	layer o	r layers is coextruded with said layer.	
24			
25	103.	A layer according to claim 99, wherein one or more of said additional	
26	layer o	r layers is laminated onto said layer.	
27			
28	104.	A layer according to claim 99, wherein one or more of said additional	
29	layer o	r layers is coated onto said layer.	

1 105. A layer according to claim 99, wherein said layer is flexible.

2

3 106. A layer according to claim 99, wherein said layer is transparent.

4

- 5 107. An article for packaging wherein the article comprises a layer
- 6 according to claim 91.

- 8 108. A process of making a polymer material by a process selected from
- 9 the group consisting of esterification, transesterification, amidation.
- 10 transamidation and direct polymerization, wherein the polymer material
- 11 comprises a polymer backbone, cyclic olefinic pendent groups, linking groups
- 12 linking the backbone with the pendent groups.
- 13 109. The process of claim 108, wherein making the polymer material
- 14 comprises the steps of:
- 15 a) selecting polymers from the group consisting of styrene/maleic
- anhydride, ethylene/maleic anhydride, ethylene/acrylic acid,
- 17 ethylene/methacrylic acid, acrylic acid, methacrylic acid, styrene/methacrylic
- 18 acid, ethylene/methyl acrylate, ethylene/ethyl acrylate, ethylene/butyl acrylate,
- 19 methyl methacrylate, methyl acrylate, and styrene/methyl methacrylate to
- 20 form a mixture and combining the polymers with a esterifying/transesterifying
- 21 compound selected from the group consisting of cyclohexene-4-methanol,
- 22 1-methyl cyclohexene-4-methanol, 2-methyl cyclohexene-4-methanol.
- 23 5-methyl cyclohexene-4-methanol, 1,2-dimethyl cyclohexene-4-methanol,
- 24 1,5-dimethyl cyclohexene-4-methanol, 2,5-dimethyl cyclohexene-4-methanol,
- 25 1,2,5-trimethyl cyclohexene-4-methanol, cyclohexene-4-ethanol, 1-methyl
- 26 cyclohexene-4-ethanol, 2-methyl cyclohexene-4-ethanol, 5-methyl
- 27 cyclohexene-4-ethanol, 1,2-dimethyl cyclohexene-4-ethanol, 1,5-dimethyl
- 28 cyclohexene-4-ethanol, 2,5-dimethyl cyclohexene-4-ethanol, 1,2,5-trimethyl

- 1 cyclohexene-4-ethanol, cyclohexene-4-propanol, 1-methyl cyclohexene-4-
- 2 propanol, 2-methyl cyclohexene-4-propanol, 5-methyl cyclohexene-4-
- 3 propanol, 1,2-dimethyl cyclohexene-4-propanol, 1,5-dimethyl cyclohexene-4-
- 4 propanol, 2,5-dimethyl cyclohexene-4-propanol, 1,2,5-trimethyl cyclohexene-
- 5 4-propanol, cyclopentene-4-methanol, 1-methyl cyclopentene-4-methanol, 3-
- 6 methyl cyclopentene-4-methanol, 1,2-dimethyl cyclopentene-4-methanol, 3,5-
- 7 dimethyl cyclopentene-4-methanol, 1,3-dimethyl cyclopentene-4-methanol,
- 8 2,3-dimethyl cyclopentene-4-methanol, 1,2,3-trimethyl cyclopentene-4-
- 9 methanol, 1,2,3,5-tetramethyl cyclopentene-4-methanol, cyclopentene-4-
- 10 ethanol, 1-methyl cyclopentene-4-ethanol, 3-methyl cyclopentene-4-ethanol,
- 11 1,2-dimethyl cyclopentene-4-ethanol, 3,5-dimethyl cyclopentene-4-ethanol,
- 12 1,3-dimethyl cyclopentene-4-ethanol, 2,3-dimethyl cyclopentene-4-ethanol,
- 13 1,2,3-trimethyl cyclopentene-4-ethanol, 1,2,3,5-tetramethyl cyclopentene-4-
- ethanol, cyclopentene-4-propanol, 1-methyl cyclopentene-4-propanol, 3-
- methyl cyclopentene-4-propanol, 1,2-dimethyl cyclopentene-4-propanol, 3,5-
- dimethyl cyclopentene-4-propanol, 1,3-dimethyl cyclopentene-4-propanol,
- 17 2,3-dimethyl cyclopentene-4-propanol, 1,2,3-trimethyl cyclopentene-4-
- propanol, and 1,2,3,5-tetramethyl cyclopentene-4-propanol;
- 19 b) heating the polymers and esterifying/transesterifying compounds
- 20 selected in (a) to form a polymer melt;
- 21 c) processing the melt in an extruder under esterification/transesterification
- 22 conditions with esterification/transesterification catalysts and antioxidants
- 23 protecting the melt from oxidation during extrusion, so that the polymer melt
- 24 undergoes esterification of polymeric anhydrides with cyclic olefin pendent
- 25 groups, esterification of polymeric acids with cyclic olefin pendent groups or
- 26 exchange of alkyl groups of polymeric esters with cyclic olefin pendent
- 27 groups; and
- 28 d) removing volatile organic products and by-products from the melt.

- 1 110. The process of claim 108, wherein making the polymer material
- 2 comprises the steps of:
- 3 a) selecting polymers from the group consisting of styrene/maleic
- 4 anhydride, ethylene/maleic anhydride, ethylene/acrylic acid,
- 5 ethylene/methacrylic acid, acrylic acid, methacrylic acid, styrene/methacrylic
- 6 acid, ethylene/methyl acrylate, ethylene/ethyl acrylate, ethylene/butyl acrylate,
- 7 methyl methacrylate, methyl acrylate, and styrene/methyl methacrylate to
- 8 form a mixture and combining the polymers with a amidizing/transamidizing
- 9 compound selected from the group consisting of cyclohexene-4-methyl
- amine, 1-methyl cyclohexene-4-methyl amine, 2-methyl cyclohexene-4-methyl
- amine, 5-methyl cyclohexene-4-methyl amine, 1,2-dimethyl cyclohexene-4-
- methyl amine, 1,5-dimethyl cyclohexene-4-methyl amine, 2,5-dimethyl
- 13 cyclohexene-4-methyl amine, 1,2,5-trimethyl cyclohexene-4-methyl amine,
- 14 cyclohexene-4-ethyl amine, 1-methyl cyclohexene-4-ethyl amine, 2-methyl
- 15 cyclohexene-4-ethyl amine, 5-methyl cyclohexene-4-ethyl amine, 1,2-dimethyl
- 16 cyclohexene-4-ethyl amine, 1,5-dimethyl cyclohexene-4-ethyl amine, 2,5-
- 17 dimethyl cyclohexene-4-ethyl amine, 1,2,5-trimethyl cyclohexene-4-ethyl
- 18 amine, cyclohexene-4-propyl amine, 1-methyl cyclohexene-4-propyl amine, 2-
- methyl cyclohexene-4-propyl amine, 5-methyl cyclohexene-4-propyl amine,
- 20 1,2-dimethyl cyclohexene-4-propyl amine, 1,5-dimethyl cyclohexene-4-propyl
- 21 amine, 2,5-dimethyl cyclohexene-4-propyl amine, 1,2,5-trimethyl
- 22 cyclohexene-4-propyl amine, cyclopentene-4-methyl amine, 1-methyl
- 23 cyclopentene-4-methyl amine, 3-methyl cyclopentene-4-methyl amine, 1,2-
- 24 dimethyl cyclopentene-4-methyl amine, 3,5-dimethyl cyclopentene-4-methyl
- amine, 1,3-dimethyl cyclopentene-4-methyl amine, 2,3-dimethyl
- 26 cyclopentene-4-methyl amine, 1,2,3-trimethyl cyclopentene-4-methyl amine,
- 27 1,2,3,5-tetramethyl cyclopentene-4-methyl amine, cyclopentene-4-ethyl
- amine, 1-methyl cyclopentene-4-ethyl amine, 3-methyl cyclopentene-4-ethyl
- 29 amine, 1,2-dimethyl cyclopentene-4-ethyl amine, 3,5-dimethyl cyclopentene-

- 1 4-ethyl amine, 1,3-dimethyl cyclopentene-4-ethyl amine, 2,3-dimethyl
- 2 cyclopentene-4-ethyl amine, 1,2,3-trimethyl cyclopentene-4-ethyl amine,
- 3 1,2,3,5-tetramethyl cyclopentene-4-ethyl amine, cyclopentene-4-propyl
- 4 amine, 1-methyl cyclopentene-4-propyl amine, 3-methyl cyclopentene-4-
- 5 propyl amine, 1,2-dimethyl cyclopentene-4-propyl amine, 3,5-dimethyl
- 6 cyclopentene-4-propyl amine, 1,3-dimethyl cyclopentene-4-propyl amine, 2,3-
- 7 dimethyl cyclopentene-4-propyl amine, 1,2,3-trimethyl cyclopentene-4-propyl
- 8 amine, and 1,2,3,5-tetramethyl cyclopentene-4-propyl amine;
- 9 b) heating the polymers and amidizing/transamidizing compounds selected
- in (a) to form a polymer melt;
- 11 c) processing the melt in an extruder under amidation/transamidation
- 12 conditions with amidation/transamidation catalysts and antioxidants protecting
- 13 the melt from oxidation during extrusion, so that the polymer melt undergoes
- 14 amidation of polymeric anhydrides with cyclic olefin pendent groups.
- 15 amidation of polymeric acids with cyclic olefin pendent groups or exchange of
- 16 alkyl groups of polymeric esters with cyclic olefin pendent groups; and
- 17 d) removing volatile organic products and by-products from the melt.
- 18 111. The process of claim 108, wherein the making of the polymer material
- 19 comprises the steps of:
- 20 (a) adding to an autoclave, ethylene and a vinyl monomer comprising 21 a pendent cyclohexene;
- 22 (b) stirring the ethylene and the vinyl monomer in the autoclave to achieve a mixture:
- 24 (c) adding a polymerization initiator before, during or after the stirring step;
- 26 (d) polymerizing the mixture to achieve a polymer; and
- (e) isolating and purifying the polymer.

112. The process of claim 111, wherein in step (a) an alpha-olefin is added to
 the autoclave along with the ethylene and the vinyl monomer and in
 step (b) the alpha-olefin is stirred with the ethylene and the vinyl
 monomer to achieve the mixture.

- 7 113. The process of claim 109, wherein the polymeric backbone is ethylenic
- 8 and the linking groups are selected from the group consisting of:

- where R is hydrogen or an alkyl group selected from the group
- consisting of methyl, ethyl, propyl and butyl groups and where n is an
- integer in the range from 1 to 12.
- 15 114. The process of claim 110, wherein the polymeric backbone is ethylenic
- 16 backbone and the linking group is:

$$-(C=O)-NH-(CHR)_n$$

- where R is hydrogen or an alkyl group selected from the group
- consisting of methyl, ethyl, propyl and butyl groups and where n is an
- integer in the range from 1 to 12.
- 21 115. The process of claim 108, wherein the material is an oxygen scavenging
- 22 composition further comprising a transition metal catalyst.

2	metal salt.
4 5	117. The process of claim 116, wherein the metal in the metal salt is cobalt.
6	118. The process according to claim 116, wherein the metal salt is selected
7	from the group consisting of cobalt neodecanoate, cobalt 2-ethylhexanoate,
8	cobalt oleate and cobalt stearate.
9	
10	
11	119. The process of claim 115, wherein the oxygen scavenging composition
12	further comprises at least one triggering material to enhance initiation of
13	oxygen scavenging.
14	
15	120. The process of claim 119, wherein the triggering material is a
16	photoinitiator.
17	
18	
19	121. The process of claim 108, wherein the cyclic olefinic pendent groups
20	have the structure (II):

 $\begin{array}{c} 1\\ 2\\ 3\\ 4\\ \end{array}$

5

- where q₁, q₂, q₃, q₄, and r are selected from the group consisting of -H,

 -CH₃, and -C₂H₅; and where m is -(CH₂)_n- and where n is an integer in

 the range of from 0 to 4; and wherein when r is -H, at least one of q₁, q₂,

 q₃ and q₄ is -H.
- 10 122. The process of claim 108, wherein the functional groups with attached 11 cyclic olefinic pendent groups are grafted onto the linking backbone by a 12 esterification, transesterification, amidation or transamidation reaction.

- 14 123. The process of claim 108, wherein the reaction is a solution reaction ora reactive extrusion.
- 16 124. The process of claim 108, wherein the esterification, transesterification,
- 17 amidation or transamidation reaction is catalyzed by a catalyst selected from
- 18 the group consisting of strong non-oxidizing acids, tertiary amines, Group I
- 19 alkoxides, Group IVB alkoxides, Group IVA organometallics.
- 20 125. The process of claim 124, wherein the catalyst is selected from the
- 21 group consisting of toluene sulfonic acid, sodium methoxide, tetrabutyl
- 22 titanate, tetraisopropyl titanate, tetra-n-propyl-titanate, tetraethyl titanate, 2-
- 23 hydroxy-pyridine and dibutyltin dilaurate.

13

15

- 1 126. The process of claim 108, wherein the backbone, linking groups and
- 2 cyclic olefin pendent groups comprise repeating units, each unit having a
- 3 structure (III) as follows:

4
5
6
$$q_3$$
 q_4
 q_2
 q_3
 q_4
 q_2
 q_3
 q_4
 q_2

- wherein P + T + Q is 100 mol % of the total composition; P is greater than 0;
- Z is selected from the group consisting of an aryl group, -(C=O)OR₁,
- 12 -O(C=O)R₁ and an alkyl aryl group, structure (IV):

14 (IV)

where R₄ is selected from the group consisting of –H, -CH₃ and -C₂H₅; R₁ is selected from the group consisting of -H, -CH₃, -C₂H₅, -C₃H₅ and -C₄H₁; R₂ and R₃ is selected from the group consisting of -H and CH₃; X is selected from the group consisting of O-, -NH-, -(C=O)O-, -(C=O)NH-, -(C=O)S-, -O(C=O)- and -(CHR)₂-; ℓ is an integer selected from the group consisting of 1, 2, 3, 4, 5 and 6; Y is -(CHR)ₙ-, where n is an integer in the range from 0 to 12 where R is selected from the group

'	consisting of $-n$, $-cn_3$ and $-c_2n_5$, where q_1 , q_2 , q_3 , q_4 , and r are selected
2	from the group consisting of -H, -CH ₃ , and -C ₂ H ₅ ; and where m is
3	-(CH ₂) _n - and where n is an integer in the range of from 0 to 4; and
4	wherein when r is –H, at least one of q_1 , q_2 , q_3 and q_4 is -H.
5	127. The process of claim 108, wherein the cyclic olefinic pendent groups are
6	selected from the group consisting of cyclohexene-4-methylene radical,
7	1-methyl cyclohexene-4-methylene radical, 2-methyl cyclohexene-4-
8	methylene radical, 5-methyl cyclohexene-4-methylene radical,
9	1,2-dimethyl cyclohexene-4-methylene radical, 1,5-dimethyl
10	cyclohexene-4-methylene radical, 2,5-dimethyl cyclohexene-4-
11	methylene radical, 1,2,5-trimethyl cyclohexene-4-methylene radical,
12	cyclohexene-4-ethylene radical, 1-methyl cyclohexene-4-ethylene
13	radical, 2-methyl cyclohexene-4-ethylene radical, 5-methyl
14	cyclohexene-4-ethylene radical, 1,2-dimethyl cyclohexene-4-ethylene
15	radical, 1,5-dimethyl cyclohexene-4-ethylene radical, 2,5-dimethyl
16	cyclohexene-4-ethylene radical, 1,2,5-trimethyl cyclohexene-4-ethylene
17	radical, cyclohexene-4-propylene radical, 1-methyl cyclohexene-4-
18	propylene radical, 2-methyl cyclohexene-4-propylene radical, 5-methyl
19	cyclohexene-4-propylene radical, 1,2-dimethyl cyclohexene-4-propylene
20	radical, 1,5-dimethyl cyclohexene-4-propylene radical, 2,5-dimethyl
21	cyclohexene-4-propylene radical, 1,2,5-trimethyl cyclohexene-4-
22	propylene radical, cyclopentene-4-methylene radical, 1-methyl
23	cyclopentene-4-methylene radical, 3-methyl cyclopentene-4-methylene
24	radical, 1,2-dimethyl cyclopentene-4-methylene radical, 3,5-dimethyl
25	cyclopentene-4-methylene radical, 1,3-dimethyl cyclopentene-4-
26	methylene radical, 2,3-dimethyl cyclopentene-4-methylene radical,
27	1,2,3-trimethyl cyclopentene-4-methylene radical, 1,2,3,5-tetramethyl
28	cyclopentene-4-methylene radical, cyclopentene-4-ethylene radical,
29	1-methyl cyclopentene-4-ethylene radical, 3-methyl cyclopentene-4-

7		etnylene radical, 1,2-dimetnyl cyclopentene-4-etnylene radical,
2		3,5-dimethyl cyclopentene-4-ethylene radical, 1,3-dimethyl
3		cyclopentene-4-ethylene radical, 2,3-dimethyl cyclopentene-4-ethylene
4		radical, 1,2,3-trimethyl cyclopentene-4-ethylene radical,
5		1,2,3,5-tetramethyl cyclopentene-4-ethylene radical, cyclopentene-4-
6		propylene radical, 1-methyl cyclopentene-4-propylene radical, 3-methyl
7		cyclopentene-4-propylene radical, 1,2-dimethyl cyclopentene-4-
8		propylene radical, 3,5-dimethyl cyclopentene-4-propylene radical,
9		1,3-dimethyl cyclopentene-4-propylene radical, 2,3-dimethyl
10		cyclopentene-4-propylene radical, 1,2,3-trimethyl cyclopentene-4-
11		propylene radical, and 1,2,3,5-tetramethyl cyclopentene-4-propylene
12		radical.
13	128.	The process of claim 108, wherein the composition is a ethylene/methyl
14		acrylate/cyclohexenyl methyl acrylate terpolymer, a cyclohexenyl methyl
15		acrylate/ethylene copolymer, a cyclohexenyl methyl
16		methacrylate/styrene copolymer, a cyclohexenyl methyl acrylate
17		homopolymer or a methyl acrylate/cyclohexenyl methyl acrylate
18		copolymer.

129.A non-odorous oxygen scavenging polymer composition comprising: (1)
 monomers derived from cyclic hydrocarbon moieties having at least one
 cyclic allylic or cyclic benzylic hydrogen and (2) a transition metal

6 oxidation catalyst.

 130. The composition of claim 129, wherein the composition comprises condensation polymers selected from the group consisting of polyesters, polyamides, polycarbonate, polyurethane, polyureas and polyether.

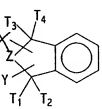
131. The composition of claim 129, wherein the cyclic allylic monomers are selected from the group consisting of structure (V), structure (VI) and structure (VII):

1 with K, L, T_1 , T_2 , T_3 , and T_4 being selected from the group consisting of $\mbox{-}\mbox{C}_{q}\mbox{H}_{\mbox{\scriptsize 2q+1}}$ with q being an integer in the range from 0 to 12 and wherein, 2 3 when either K or L is -H, at least one of T_1 , T_2 , T_3 and T_4 is -H; 4 and with X and Y being selected from the group consisting of -(CH₂)_n-OH, -(CH₂)_n-NH₂, -(CH₂)_nNC=O and -(CH₂)_m-(C=O)-A with n being an 5 6 integer in the range from 1 to 12 and m being an integer in the range 7 from 0 to 12 and A being selected from the group consisting of --OH, -OCH₃, -OC₂H₅, -OC₃H₇ and halides; and Q being selected from the 8 9 group consisting of -(CtH2t-2) with t being an integer in the range from 1 10 to 4; and with G being selected from –(C=O)- and –(C_nH_{2n+1})- with n being an 11 12 integer from 0 to 12. 13 14 132. The composition of claim 129, wherein the cyclic benzylic monomers are 15 selected from the group consisting of structure (VIII), structure (IX), structure 16 (X), structure (XI), structure (XII), and structure (XIII) 17 18 (VIII) (IX) 19

18 (VIII)
19 T₃ T₄
20
21 Z
7 T₁ T₂

232425

2627



(X)

$$Z = \begin{bmatrix} T_4 & G \\ & & \\ T_1 & T_2 & 0 \end{bmatrix}$$

$$T_3$$
 Z
 CH_2
 CH_2

(XII)

$$0 = \begin{bmatrix} T_1 & T_2 \\ 0 & T_3 & T_4 \end{bmatrix}$$

(XIII)
$$\hat{O}$$

$$CH_2$$

$$T_1$$

$$T_2$$

$$CH_2$$

$$T_3$$

$$T_4$$

1	where X and Y are selected from the group consisting of –(CH ₂) _n -OH,
2	-(CH ₂) _n -NH ₂ and -(CH ₂) _m -(C=O)-R ₁ with n being an integer in the range
3	from 1 to 12, and with m being an integer in the range from 0 to 12 and
4	with R₁ being selected from the group consisting of –OH, -OCH₃,
5	-OC ₂ H ₅ , -OC ₃ H ₇ and halides;
6	with T_1 , T_2 , T_3 , and T_4 being selected from the group consisting of
7	$-C_qH_{2q+1}$ with q being an integer in the range from 0 to 12 and at least
8	one of T_1 , T_2 , T_3 and T_4 being $-H$;
9	and with X and Y being selected from the group consisting of
10	$(CH_2)_n$ -OH, - $(CH_2)_n$ -NH ₂ , - $(CH_2)_n$ NC=O, and - $(CH_2)_m$ - $(C=O)$ -A with n
11	being an integer in the range from 1 to 12, and m being an integer in the
12	range from 0 to 12 and A being selected from the group consisting of
13	-OH, -OCH ₃ , -OC ₂ H ₅ , -OC ₃ H ₇ and halides; and Z being selected from the
14	group consisting of -(C_tH_{2t-2})-, -O-, -NR ₂ -, -S-, with t being an integer in
15	the range from 1 to 4 and R ₂ being selected from the group consisting of
16	–OH, -OCH ₃ , -OC ₂ H ₅ , -OC ₃ H ₇ and halides;
17	and with G being selected from –(C=O)- and -(C_nH_{2n+1})- with n being an
8	integer from 0 to 12.
19	
20	133. The composition of claim 130, the composition being thermoplastic.
21	
22	134. The composition of claim 130, the composition being thermoset.
23	
24	
25	135. The composition of claim 130, the composition being a multilayered
26	structure with other layers being an aromatic polyester or copolyester
27	selected from the group consisting of polyethylene terephthalate,
28	polyethylene naphthalate, polypropylene terephthalate, polybutylene
29	terephthalate, polyethylene isophthalate, polycyclohexanedimethanol

1	terephthalate, polybutylene naphthalate, polycyclohexanedimethanol
2	naphthalate, and copolymers and blends thereof.
3	
4	136. The composition of claim 130, the composition being a multilayered
5	structure with other layers being polyamides or copolyamides selected from
6	the group consisting of Nylon 6, Nylon 66, Nylon 610 and mixtures thereof.
7	
8	
9	137. The composition of claim 130, the composition being a multilayered
10	structure with other layers being bisphenol A carbonate.
11	
12	138. The composition of claim 130, the composition being a multilayered
13	structure with other layers being vinylic polymers or copolymers selected
14	from the group consisting of ethylene polymer, ethylene copolymer, propylene
15	polymer, propylene copolymer, styrene polymer, styrene copolymer, acrylate
16	polymer, acrylate copolymer, vinyl chloride polymer, vinyl chloride copolymer,
17	divinyl chloride polymer, divinyl chloride copolymer, fluorinated vinyl polymer,
18	fluorinated vinyl copolymer and mixtures thereof.
19	
20	
21	139. The composition of claim 130, the composition being blended with an
22	aromatic polyester or copolyester selected from the group consisting of
23	polyethylene terephthalate, polyethylene naphthalate, polypropylene
24	terephthalate, polybutylene terephthalate, polyethylene isophthalate,
25	polycyclohexandedimethanol terephthalate, polybutylene naphthalate,
26	polycyclohexanedimethanol naphthalate, and copolymers and blends thereof.
27	
28	140. The composition of claim 130, the composition being blended with
29	polyamides or copolyamides selected from the group consisting of Nylon 6,
30	Nylon 66, Nylon 610 and mixtures thereof.

1 141. The composition of claim 130, the composition being blended with 2 bisphenol A polycarbonate. 3 4 142. The composition of claim 130, the composition being a blend comprising vinvlic polymers or copolymers selected from the group consisting of ethylene 5 6 polymer, ethylene copolymer, propylene polymer, propylene copolymer. 7 styrene polymer, styrene copolymer, acrylate polymer, acrylate copolymer. vinyl chloride polymer, vinyl chloride copolymer, divinyl chloride polymer. 8 9 divinyl chloride copolymer, fluorinated vinyl polymer, fluorinated vinyl 10 copolymer and mixtures thereof. 11 12 13 143. The composition of claim 130, the composition being laminated or 14 adhering onto a substrate selected from the group consisting of paper, foil, 15 high temperature film, metallized film, polyamide films, ethylene vinyl alcohol 16 film, silica coated film, nylon/EVOH/nylon, oriented polypropylene, polyester 17 film, polyethylene, polypropylene, polyester, oriented polyethylene 18 terephthalate and cellophane. 19 20 144. The composition of claim 129, wherein the composition comprises a 21 vinyl polymer selected from the group consisting of ethylene polymer. 22 ethylene copolymer, propylene polymer, propylene copolymer, styrene 23 polymer, styrene copolymer and mixtures thereof. 24 25 26 145. A rigid container for food or beverage, the container being molded from 27 a resin comprising a non-odorous oxygen scavenging polymer composition. 28 the composition comprising (1) monomers derived from cyclic hydrocarbon 29 moieties having at least one cyclic allylic or cyclic benzylic hydrogen and (2) a 30 transition metal oxidation catalyst.

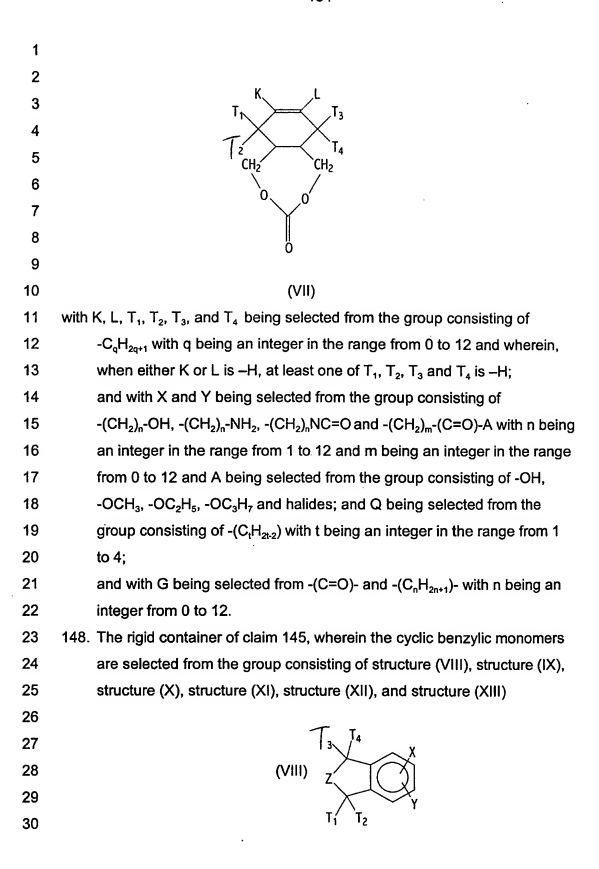
(V)

- 1 146. The rigid container of claim 145, wherein the composition comprises
- 2 condensation polymers selected from the group consisting of polyesters,
- 3 polyamides, polycarbonate, polyurethane, polyureas, polysulfones and
- 4 polyether.

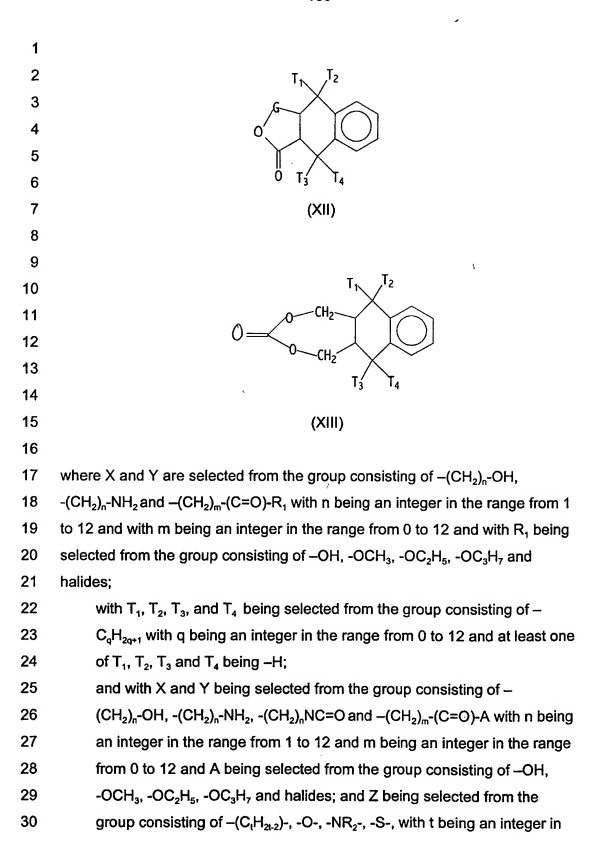
147. The rigid container of claim 145, wherein the cyclic allylic monomers are selected from the group consisting of structure (V), structure (VI) and structure (VII):

(VI)

-



1	
2	
2	
4	
5	
6	,
7	T_3
8	
9	Y
10	T_1 T_2
11	
12	(IX)
13	
14	
15	
16	
17	
18	T_3 J_4
19	Z, G
20	
21	$T_1 T_2$
22	(X)
23	
0.1	T_3 T_4 CH_2 CH
24 25	(XI)
26 26	(^1)
20	



1	the range from 1 to 4 and R ₂ being selected from the group consisting of		
2	–OH, -OCH₃, -OC₂H₅, -OC₃H₂ and halides;		
3	and with G being selected from –(C=O)- and –(C_nH_{2n+1})- with n being an		
4	integer from 0 to 12.		
5			
6	149. The rigid container of claim 146, the composition being thermoplastic.		
7	150. The rigid container of claim 146, the composition being thermoset.		
8			
9			
10	151. The rigid container of claim 146, the composition being made by		
11	coextrusion, blow molding or lamination with an aromatic polyester or		
12	copolyester selected from the group consisting of polyethylene terephthalate,		
13	polyethylene naphthalate, polypropylene terephthalate, polybutylene		
14	terephthalate, polyethylene isophthalate, polycyclohexanedimethanol		
15	terephthalate, polybutylene naphthalate, polycyclohexanedimethanol		
16	naphthalate, and copolymers and blends thereof.		
17			
18	152. The rigid container of claim 146, the composition being made by		
19	coextrusion, blow molding or lamination with polyamides or copolyamides		
20	selected from the group consisting of Nylon 6, Nylon 66, Nylon 610 and		
21	mixtures thereof.		
22			
23			
24	153. The rigid container of claim 146, the composition being made by		
25	coextrusion, blow molding or lamination with bisphenol A polycarbonate.		
26			
27	154. The rigid container of claim 146, the composition being made by		
28	coextrusion, blow molding or lamination with vinylic polymers or copolymers		
29	selected from the group consisting of ethylene polymer, ethylene copolymer,		
30 .	propylene polymer, propylene copolymer, styrene polymer, styrene		

1	copolymer, acrylate polymer, acrylate copolymer, vinyl chloride polymer, vinyl		
2	chloride copolymer, divinyl chloride polymer, divinyl chloride copolymer,		
3	fluorinated vinyl polymer, fluorinated vinyl copolymer and mixtures thereof.		
4			
5			
6	155. The rigid container of claim 146, the composition being blended with		
7	an aromatic polyester or copolyester selected from the group consisting of		
8	polyethylene terephthalate, polyethylene naphthalate, polypropylene		
9	terephthalate, polybutylene terephthalate, polyethylene isophthalate,		
10	polycyclohexandedimethanol terephthalate, polybutylene naphthalate,		
11	polycyclohexanedimethanol naphthalate, and copolymers and blends thereof.		
12			
13	156. The rigid container of claim 146, the composition being blended with		
14	polyamides or copolyamides selected from the group consisting of Nylon 6,		
15	Nylon 66, Nylon 610 and mixtures thereof.		
16			
17			
18	157. The rigid container of claim 146, the composition being blended with		
19	bisphenol A carbonate.		
20	·		
21	158. The rigid container of claim 146, the composition being blended with		
22	vinylic polymers or copolymers selected from the group consisting of ethylene		
23	polymer, ethylene copolymer, propylene polymer, propylene copolymer,		
24	styrene polymer, styrene copolymer, acrylate polymer, acrylate copolymer,		
25	vinyl chloride polymer, vinyl chloride copolymer, divinyl chloride polymer,		
26	divinyl chloride copolymer, fluorinated vinyl polymer, fluorinated vinyl		
27	copolymer and mixtures thereof.		
28			
29			

- 1 159. The rigid container of claim 146, the composition being laminated or adhering onto a substrate selected from the group consisting of paper, foil,
- 3 high temperature film, metallized film, polyamide films, ethylene vinyl alcohol
- 4 film, silica coated film, nylon/EVOH/nylon, oriented polypropylene, polyester
- 5 film, oriented polyethylene terephthalate, polypropylene, polyester, and
- 6 cellophane.
- 7 160. The rigid container of claim 146, wherein the composition comprises
- 8 a vinyl polymer selected from the group consisting of ethylene polymer,
- 9 ethylene copolymer, propylene polymer, propylene copolymer, styrene
- 10 polymer, styrene copolymer and mixtures thereof.

- 12 161. The rigid container according to claim 145, wherein the composition is
- 13 a single layer.

1415

- 16 162. The rigid container according to claim 145, wherein the composition is
- 17 multilayered.

18

- 19 163. The rigid container according to claim 162, wherein the composition
- 20 comprises an outer air contact layer and an inner oxygen scavenging layer.

21

22

- 23 164. The rigid container according to claim 163, wherein the outer air
- 24 contact layer comprises an oxygen barrier resin selected from the group
- 25 consisting of polyethylene terephthalate, polyethylene naphthalate and a
- 26 mixture of polyethylene terephthalate and polyethylene naphthalate.

- 28 165. The rigid container according to claim163, wherein the composition
- 29 further comprises at least one of an inner food contact layer, a tie layer, and a
- 30 tinted ultraviolet protection layer.

WO 99/48963

1		
2		
3	166.	The rigid container according to claim 165, wherein the inner food
4	contact	layer comprises an oxygen barrier resin selected from the group
5	consist	ing of polyethylene terephthalate, polyethylene naphthalate and a
6	mixture	of polyethylene terephthalate and polyethylene naphthalate.
7		
8	167.	The rigid container of claim 145, wherein oxygen scavenging by the
9	non-od	orous oxygen scavenging polymer composition is initiated by moisture
10	or actin	ic radiation.
11		
12		
13	168.	The rigid container of claim 145, wherein the transition metal catalyst
14	is a me	tal salt.
15		
16	169.	The rigid container of claim 168, wherein the metal in the metal salt is
17	cobalt.	
18		
19		
20	170.	The rigid container of claim 169, wherein the metal salt is selected
21	from th	e group consisting of cobalt neodecanoate, cobalt 2-ethylhexanoate,
22	cobalt	pleate and cobalt stearate.
23		
24	171.	The rigid container of claim 145, wherein the non-odorous oxygen
25	scaven	ging composition further comprises at least one triggering material to
26	enhand	e initiation of oxygen scavenging.
27		
28		
29	172.	The rigid container of claim 171, wherein the triggering material is a
30	photoin	nitiator.

4	
ı	

2 173. The rigid container of claim 172, wherein the photoinitiator has an

- 3 ultraviolet absorption window above 320 nm.
- 4 174. The rigid container of claim 172, wherein the container further comprises.
- 5 a tinted ultraviolet protection layer located between the layer comprising the
- 6 non-odorous oxygen scavenging composition and the inside of the rigid
- 7 container.

8

9 175. The rigid container of claim 174, wherein the tinted layer is the food

10 contact layer.

11

12

- 13 176. The rigid container of claim 174, wherein the resin comprises
- 14 condensation polymers selected from the group consisting of polyesters,
- 15 polyamides, polyurethane, polyureas, polysulfones, polycarbonates and
- 16 polyether.

17

- 18 177. The rigid container of claim 174, wherein the cyclic allylic monomers
- 19 are selected from the group consisting of structure (V), structure (VI) and
- 20 structure (VII):

21

22

23

24

25

25

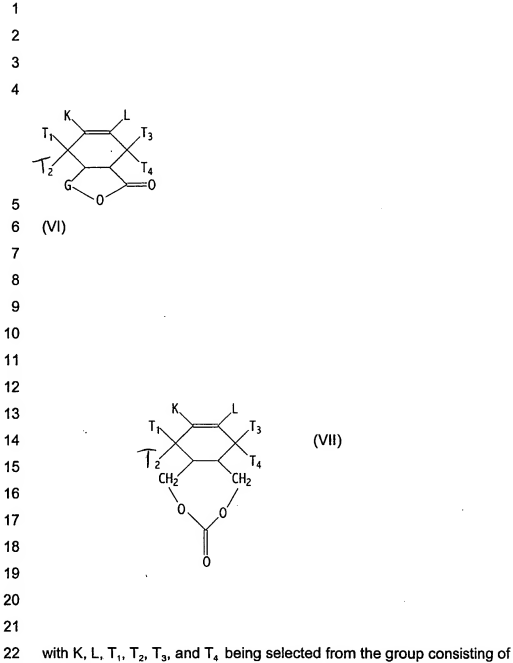
26

27

28

29

$$\uparrow_{4}
\downarrow_{Q}
\downarrow_{T_{1}}
\downarrow_{Q}
\downarrow_{T_{1}}$$
(V)



-C_qH_{2q+1} with q being an integer in the range from 0 to 12 and wherein, when either K or L is -H, at least one of T₁, T₂, T₃ and T₄ is -H; and with X and Y being selected from the group consisting of -(CH₂)_n-OH, -(CH₂)_n-NH₂, -(CH₂)_nNC=O and -(CH₂)_m-(C=O)-A with n being an integer in the range from 1 to 12 and m being an integer in the range

1	from 0 to 12 and A being s	selected from the group consisting of -OH,
2	-OCH ₃ , -OC ₂ H ₅ , -OC ₃ H ₇ ar	nd halides; and Q being selected from the
3	group consisting of -(CtH2t	₋₂) with t being an integer in the range from 1
4	to 4;	
5	and with G being selected	from -(C=O)- and -(C _n H _{2n+1})- with n being an
6	integer from 0 to 12.	·
7		
8	178. The rigid container of clair	m 174, wherein the cyclic benzylic monomers
9	are selected from the grou	up consisting of structure (VIII), structure (IX),
10	structure (X), structure (XI), structure (XII), and structure (XIII):
11		
12	(VIII)	. (IX)
13	Ta	T ₃
14	3 X	z
15	z ()	Y
16	T_1 T_2 Y	T_1' T_2
17	.1 .2	
18		•
19		
20		
21		
22		
23	:	
24		
25		
26		
27		
28		» J4
29	(X)	, G
30		Ž O
		T_1 T_2 0
		V

where X and Y are selected from the group consisting of -(CH₂)_n-OH,

-(CH₂)_n-NH₂ and -(CH₂)_m-(C=O)-R₁ with n being an integer in the range

from 1 to 12 and with m being an integer in the range from 0 to 12 and

WO 99/48963

1		with R_1 being selected from the group consisting of -OH, -OCH $_3$, -OC $_2$ H $_5$,
2		-OC ₃ H ₇ and halides;
3		with T_1 , T_2 , T_3 , and T_4 being selected from the group consisting of
4		$-C_qH_{2q+1}$ with q being an integer in the range from 0 to 12 and at least
5		one of T_1 , T_2 , T_3 and T_4 being -H;
6		and with X and Y being selected from the group consisting of
7		-(CH ₂) _n -OH, -(CH ₂) _n -NH ₂ , -(CH ₂) _n NC=O and -(CH ₂) _m -(C=O)-A with n
8		being an integer in the range from 1 to 12 and m being an integer in the
9		range from 0 to 12 and A being selected from the group consisting of
10		-OH, -OCH ₃ , -OC ₂ H ₅ , -OC ₃ H ₇ and halides; and Z being selected from the
11		group consisting of -(C_tH_{2t-2})-, -O-, -NR ₂ -, -S-, with t being an integer in
12		the range from 1 to 4 and R ₂ being selected from the group consisting of
13		-OH, -OCH ₃ , -OC ₂ H ₅ , -OC ₃ H ₇ and halides;
14		and with G being selected from -(C=O)- and -(C_nH_{2n+1})- with n being an
15		integer from 0 to 12.
16		
17	179.	The rigid container of claim 176, the composition being thermoplastic.
18		
19	180.	The rigid container of claim 176, the composition being thermoset.
20		·
21		
22	181.	The rigid container of claim 176, the composition being in a multilayer
23		structure with other layers, at least one of the other layers comprising an
24		aromatic polyester or copolyester selected from the group consisting of
25		polyethylene terephthalate, polyethylene naphthalate, polypropylene
26		terephthalate, polybutylene terephthalate, polyethylene isophthalate,
27		polycyclohexanedimethanol terephthalate, polybutylene naphthalate,
28		polycyclohexanedimethanol naphthalate, and copolymers and blends
29		thereof.
30		

-166-

1 182. The rigid container of claim 176, the composition being in a multilayer 2 structure with other layers, at least one of the other layers comprising 3 polyamides or copolyamides selected from the group consisting of Nylon 6, 4 Nylon 66, Nylon 610 and mixtures thereof. 5 6 183. The rigid container of claim 176, the composition being in a multilayer 7 structure with other layers, at least one of the other layers comprising 8 bisphenol A carbonate. 9 10 11 184. The rigid container of claim 176, the composition being in a multilayer 12 structure with other layers, at least one of the other layers comprising vinylic 13 polymers or copolymers selected from the group consisting of ethylene 14 polymer, ethylene copolymer, propylene polymer, propylene copolymer, 15 styrene polymer, styrene copolymer, acrylate polymer, acrylate copolymer, 16 vinyl chloride polymer, vinyl chloride copolymer, divinyl chloride polymer, 17 divinyl chloride copolymer, fluorinated vinyl polymer, fluorinated vinyl 18 copolymer and mixtures thereof. 19 . 185. The rigid container of claim 176, the composition being blended with an 20 21 aromatic polyester or copolyester selected from the group consisting of 22 polyethylene terephthalate, polyethylene naphthalate, polypropylene 23 terephthalate, polybutylene terephthalate, polyethylene isophthalate, 24 polycyclohexandedimethanol terephthalate, polybutylene naphthalate, 25 polycyclohexanedimethanol naphthalate, and copolymers and blends thereof. 26 27 28 186. The rigid container of claim 176, the composition being blended with 29 polyamides or copolyamides selected from the group consisting of Nylon 6, 30 Nylon 66, Nylon 610 and mixtures thereof.

1	
2	187. The rigid container of claim 176, the composition being blended with
3	bisphenol A carbonate.
4	
5′	
6	188. The rigid container of claim 176, the composition being blended with
7	vinylic polymers or copolymers selected from the group consisting of ethylene
8	polymer, ethylene copolymer, propylene polymer, propylene copolymer,
9	styrene polymer, styrene copolymer, acrylate polymer, acrylate copolymer,
10	vinyl chloride polymer, vinyl chloride copolymer, divinyl chloride polymer,
11	divinyl chloride copolymer, fluorinated vinyl polymer, fluorinated vinyl
12	copolymer and mixtures thereof.
13	
14	189. The rigid container of claim 176, the composition being laminated or
15	adhering onto a substrate selected from the group consisting of paper, foil,
16	high temperature film, metallized film, polyamide films, ethylene vinyl alcohol
17	film, silica coated film, nylon/EVOH/nylon, oriented polypropylene, polyester
18	film, polyethylene, polypropylene, oriented polyethylene terephthalate, and
19	cellophane.
20	
21	
22	190. The rigid container of claim 176, wherein the composition comprises a
23	vinyl polymer selected from the group consisting of ethylene polymer,
24	ethylene copolymer, propylene polymer, propylene copolymer, styrene
25	polymer, styrene copolymer and mixtures thereof.
26	191. The rigid container according to claim 176, wherein the composition
27	comprises an outer air contact layer and an inner oxygen scavenging layer.
28	
29	192. The rigid container according to claim 191, wherein the outer air contact
30	layer comprises an oxygen barrier resin selected from the group consisting of

polyethylene terephthalate, polyethylene naphthalate and a mixture of 1 2 polyethylene terephthalate and polyethylene naphthalate. 3 4 5 193. The rigid container according to claim 176, wherein the composition 6 further comprises at least one of an inner food contact layer, a tie layer, and a 7 tinted ultraviolet protection layer. 8 9 194. The rigid container according to claim 193, wherein the inner food 10 contact layer comprises an oxygen barrier resin selected from the group 11 consisting of polyethylene terephthalate, polyethylene naphthalate and a 12 mixture of polyethylene terephthalate and polyethylene naphthalate. 13 14 15 195. The rigid container of claim 176, wherein the transition metal catalyst is 16 a metal salt. 17 18 196. The rigid container of claim 195, wherein the metal in the metal salt is 19 cobalt. 20 21 22 197. The rigid container of claim 196, wherein the metal salt is selected from 23 the group consisting of cobalt neodecanoate, cobalt 2-ethylhexanoate, cobalt 24 oleate and cobalt stearate. 25 198. The rigid container of claim 197, wherein the triggering material is a 26 photoinitiator. 27 28 199. The rigid container of claim 198, wherein the photoinitiator has an 29 ultraviolet absorption window above 320 nm. 30

- 2 200. The rigid container of claim 145, wherein the rigid container is suitable
- 3 for packaging oxygen sensitive drinks for extended freshness and shelf life.

- 5 201. The rigid container of claim 200, wherein the oxygen sensitive drink is
- 6 beer.

$$CH_{3}$$

$$3-Cyclohexene-1-methanol$$

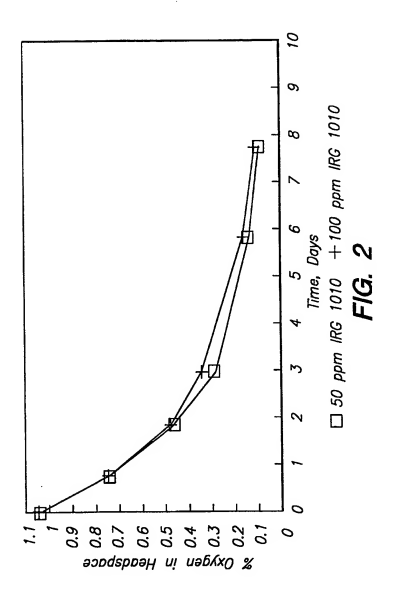
$$Reactive Extrusion (REX)$$

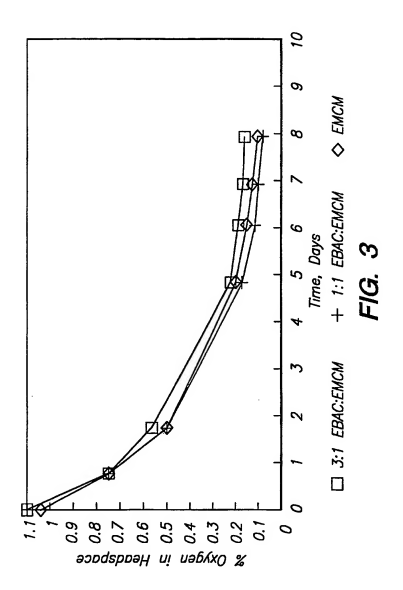
$$CH_{3}$$

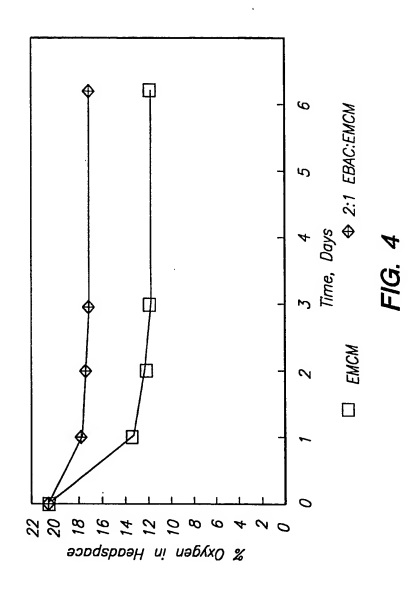
$$C=0$$

EMCM poly(ethylene/methyl acrylate/cyclohexene-methyl acrylate)

FIG. 1







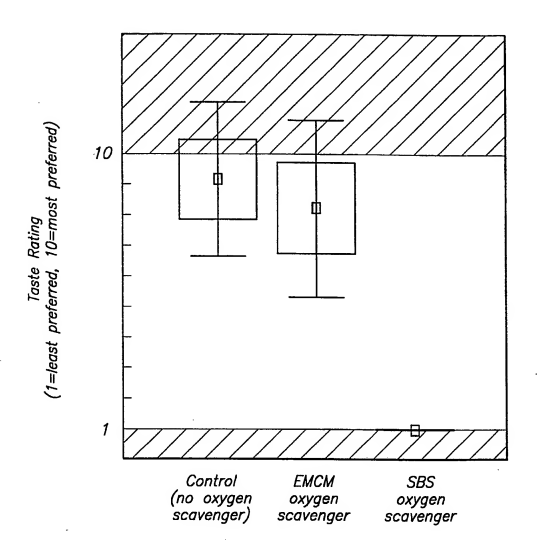


FIG. 5



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

C08K 5/09

(11) International Publication Number:

WO 99/48963

A3

(43) International Publication Date: 30 September 1999 (30.09.99)

(21) International Application Number:

PCT/US99/06379

(22) International Filing Date:

24 March 1999 (24.03.99)

(30) Priority Data:

330077 09/127,316 25 March 1998 (25.03.98) NZ

31 July 1998 (31.07.98) US

(71) Applicants: CHEVRON CHEMICAL COMPANY LLC [US/US]; 555 Market Street, San Francisco, CA 94105 (US). SEALED AIR (NZ) LIMITED [NZ/NZ]; Private Bag 3085, Corner Avalon Drive & Foreman Road (NZ).

(72) Inventors: CHING, Ta, Yen; 10 Santa Yorma Court, Novato, CA 94945 (US). CAI, Gangfeng; 335 Glasgow Circle, Danville, CA 94526 (US). DEPREE, Craig; 27 Pembroke Street, Palmerston North (NZ). GALLAND, Mark, Steven; 2806 Chasse Ridge Drive, Orange, TX 77632 (US). GOODRICH, Joseph, L.; 3545 Wilkinson Lane, Lafayette, CA 94549 (US). LEONARD, James, P.; 545 Bret Harte Road, San Rafael, CA 94901 (US). MATTHEWS, Andrew; 106 West Shallowstone Road, Greer, SC 29650 (US). RUSSELL, Kenneth, W.; Route 10, Box 1366, Orange, TX 77630 (US). YANG, Hu; 360 Bellflower Court, San Ramon, CA 94583 (US).

(74) Agents: HAYMOND, W., Bradley et al.; Chevron Corporation, Law Dept., P.O. Box 7141, San Francisco, CA 94120-7141 (US).

(81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(88) Date of publication of the international search report:

11 November 1999 (11.11.99)

(54) Title: OXYGEN SCAVENGERS WITH REDUCED OXIDATION PRODUCTS FOR USE IN PLASTIC FILMS AND BEVERAGE AND FOOD CONTAINERS

ЕМСМ poly(ethylene/methyl acrylate/cyclohexene-methyl acrylate)

(57) Abstract

An oxygen scavenger composition, for use in or with plastics materials, includes a polymer or oligomer having at least one cyclohexene group or functionality. The composition produces only low levels of volatile or (extractable from a plastics material in which it is incorporated) products as a consequence of oxygen scavenging. A family of polymers containing selected cyclic allylic pendent groups for oxygen scavenging packaging which has minimal organoleptic by-products after oxidation. Multilayer plastic containers for food and beverage packaging which incorporate oxygen scavenging resins which selectively oxidize upon activation without giving off odorous fragments.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan ·	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	lT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
Cl	Côte d'Ivoire	KР	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

International Application No PCT/US 99/06379

PCT/US 99/06379 A. CLASSIFICATION OF SUBJECT MATTER IPC 6 C08K5/09 According to International Patent Classification (IPC) or to both national classification and IPC Minimum documentation searched (classification system followed by classification symbols) IPC 6 C08K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category ° Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. Χ -WO 96 40799 A (CHEVRON CHEM CO) 1-30 19 December 1996 (1996-12-19) cited in the application page 30, line 14 - page 31, line 22; claims 1,3; example 44; tables 4,7-9 EP 0 040 983 A (UNIV CALIFORNIA) Х 1,5 2 December 1981 (1981-12-02) page 14, line 10-15; claims 1,3,7,8 DE 19 00 181 A (DEUTSCHE AKADEMIE DER Χ 1.5 WISSENSCAHFTEN BERLIN) 2 October 1969 (1969-10-02) page 3, line 20-30; example 1 EP 0 418 011 A (SERICOL GROUP LTD) Х 1,5 20 March 1991 (1991-03-20) claims 1-6; example 1 -/--Further documents are listed in the continuation of box C. , Patent family members are listed in annex. * Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application *A* document defining the general state of the art which is not considered to be of particular relevance cited to understand the principle or theory underlying the invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another involve an inventive step when the document is taken alone 'Y' document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or other means ments, such combination being obvious to a person skilled document published prior to the international filing date but later than the priority date claimed *&* document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 2 4. 0<u>9. **99**</u> 15 June 1999 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Devriese, K

International Application No PCT/US 99/06379

	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
	WO 95 02616 A (CHEVRON RES & TECH) 26 January 1995 (1995-01-26) cited in the application	1
	·	
	-	·
: :		

International application No.

PCT/US 99/06379

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
See additional sheet
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. X No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1 - 30
Remark on Protest The additional search fees were accompanied by the applicant's protest.
No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

1. Claims: 1-30

a composition comprising: a) a polymer or oligomer with cyclohexene functionality and b)a transition metal catalyst;

2. Claims: 30-107

a polymer with cyclic olefinic pendant groups, a composition and article thereof

3. Claims: 31-41,108-128

a polymer with cyclic olefinic pendant groups, a process for its manufacture

4. Claims: 129-201

a composition containing a) a polymer b) monomers having a cyclic allylic hydrogen and c) a transition metal catalyst; a container moulded from said composition.

5. Claims: 129-201

a composition containing a) a polymer b) monomers having a cyclic benzylic hydrogen and c) a transition metal catalyst; a container moulded from said composition.

Information on patent family members

International Application No PCT/US 99/06379

Patent document cited in search report		Publication date	Patent family member(s)		Publication date -
WO 9640799	A	19-12-1996	AU 563269 BR 960923 CA 221838 EP 083038	2 A 1 A	30-12-1996 11-05-1999 19-12-1996 25-03-1998
EP 0040983	A	02-12-1981	JP 5701180	1 A	21-01-1982
DE 1900181	A	02-10-1969	CS 15150	2 B	19-10-1973
EP 0418011	Α	20-03-1991	GB 223592 AT 16329 AU 610579 DE 6903205 JP 323845	8 T 0 A 5 D	20-03-1991 15-03-1998 21-03-1991 26-03-1998 24-10-1991
WO 9502616	A	26-01-1995	AU 69620 AU 375079 AU 68481 AU 736109 CA 214430 DE 6941096 EP 065919 JP 850518 NO 95092 NZ 26993 US 573663 US 585914	07 A 00 B 04 A 052 D 052 T 053 T 064 A 069 A	03-09-1998 20-11-1997 08-01-1998 13-02-1995 26-01-1995 16-07-1998 08-10-1998 28-06-1995 04-06-1996 04-05-1995 27-08-1996 07-04-1998 06-05-1997 12-01-1999